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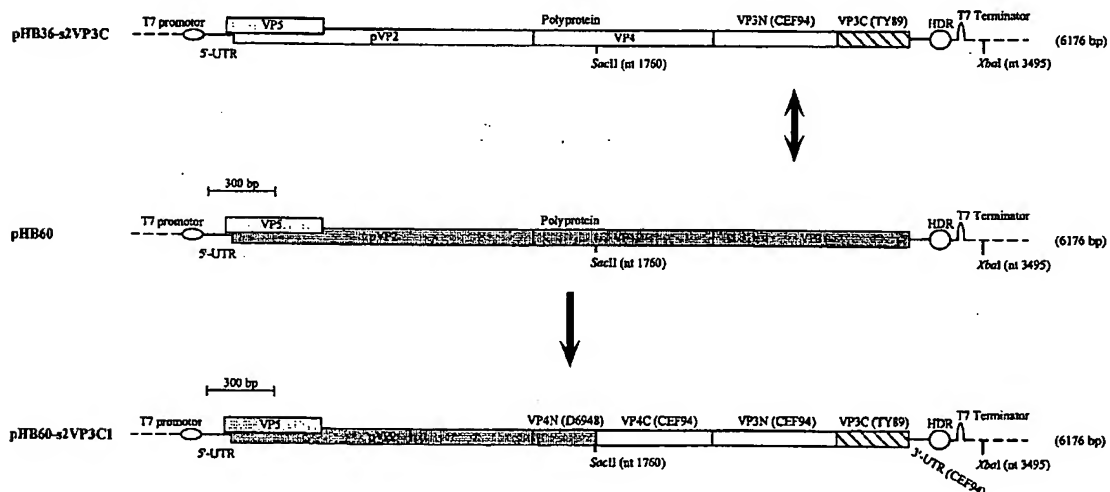
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### (54) Mosaic Infectious Bursal Disease Virus vaccines

(57) The invention relates to recombinant Infectious Bursal Disease Virus (IBDV) and vaccines derived thereof. The invention provides infectious recombinant Infectious Bursal Disease Virus (rIBDV) essentially in-

capable of growing in a cell that is not derived from a bursa cell, or an infectious rIBDV having retained at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus (vvIBDV).

Fig. 5g Schematic representation of the construction of plasmid pHB60-s2VP3C1



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**Description**

[0001] The invention relates to Infectious Bursal Disease Virus (IBDV) vaccines.

Infectious Bursal Disease (IBD), an infectious disease among young chickens, was first recognized in 1957 in Gumboro, Delaware, USA and formally documented by Cosgrove (Cosgrove, 1962; Lasher and Shane, 1994). As a result, the disease is often referred to as Gumboro. Not long after IBD was first reported, it was being recognized in poultry populations throughout the world (Lasher and Shane, 1994). IBD is caused by a virus (IBDV) classified as a Birnavirus (Dobos et al., 1979). Two different IBDV serotypes exist: serotype I and II (Jackwood et al., 1982; McFerran et al., 1980). Isolates belonging to serotype I are highly pathogenic for chickens. Serotype II isolates, which are mainly recovered from turkeys, have never been reported to induce clinical signs in chickens and are regarded as apathogenic (Ismail et al., 1988). Infectious bursal disease or Gumboro is a highly contagious disease for young chickens, and is responsible for severe losses in poultry industries. In birds surviving an acute infection, lymphoid cells in the bursa of Fabricius are destroyed, resulting in B-cell dependent immunodeficiency. This causes increased susceptibility to disease caused by otherwise harmless agents. A central role in the pathogenesis of Gumboro is played by the bursa, which is representing the target organ of the virus.

[0002] IBDV infections were initially recognized by whitish or watery diarrhea, anorexia, depression, trembling, weakness, and death. This clinical IBD was generally seen in birds between three and eight weeks of age. The course of the disease runs approximately 10 day in a flock. Mortality usually ranges from 0-30 percent. Field reports suggest that leghorns are more susceptible to IBDV than broiler type chickens. Subclinical IBD was later recognized and is generally considered a greater problem in commercial poultry than the clinical disease. It is generally seen in birds less than three weeks of age. This early infection results in a B-lymphocyte depletion of the bursa of Fabricius. The bird is immunologically crippled and unable to respond fully to vaccinations or field infections. In susceptible chickens, damage caused by IBDV can be seen within two to three days after exposure to virulent virus. Initially, the bursa swells (3 days post-exposure) with edema and hemorrhages and then begins to show atrophy (7-10 days). IBD virus is especially cytopathic to certain B-lymphocytes. The highest concentration of these specific B-lymphocytes is found in the bursa. Destruction of the B-lymphocytes by IBD field virus may result in an incomplete seeding of these cells in secondary lymphoid tissue. As a result of the depletion of B-lymphocytes, surviving birds are immunocompromised during the remaining of their live time.

[0003] IBDV is found worldwide, and IBDV specific antibodies have even been found in Antarctic penguins (Gardner et al., 1997). The prevalence of clinical IBD is relatively low compared to the prevalence of subclinical IBD. IBDV is very resistant to common disinfectants and has been found in lesser mealworms, mites, and mosquitoes. These facts correlate with field experience of reoccurring IBD problems on a farm, despite clean-up efforts. Infection with IBDV results in a strong antibody response against IBD, which is capable of neutralizing this virus. Most likely as a result of vaccination, antigenic variant isolates of serotype I were isolated in the Delaware area (USA). These isolates have been shown to cause bursa atrophy in as little as three days post-infection without inflammation of the bursa. Despite their change in antigenicity these antigenic variants do not form a distinct serotype. After the occurrence of antigenic variant IBDV isolates in the USA, the poultry industry in European countries was hit by outbreaks of IBD caused by a very virulent serotype I IBDV (vvIBDV) (Berg et al., 1991; Chettle et al., 1989; Kouwenhoven and Van den Bos, 1995). These very virulent field isolates were capable of establishing themselves in the face of high levels of maternal antibodies which normally were protective. These vvIBDV cause more severe clinical signs during an outbreak and are now found globally (e.g. Europe, Japan, Israel and Asia).

[0004] IBDV belongs to the family of Birna viruses which include Infectious Bursal Disease Virus (IBDV) isolated from chickens, Infectious Pancreatic Necrosis Virus (IPNV) isolated from Fish, *Drosophila* X Virus (DXV) isolated from fruit fly, and *Tellina* virus (TV) and Oyster Virus (OV) both isolated from bivalve molluscs (Dobos et al., 1979). Birna viruses have a dsRNA genome which is divided over two genome segments (the A- and B-segment). The A-segment (3.3 kbp) contains two partly overlapping open reading frames (ORFs). The first, smallest ORF encodes the non-structural Viral Protein 5 (VP5, 17 kDa). The second ORF encodes a polyprotein (1012 amino acid, 110 kDa), which is autocatalytically cleaved. The exact position of these cleavage sites is unknown. From SDS-Page analysis of *in vitro* translated IBDV RNA it is known that the polyprotein is rapidly cleaved into three proteins: pVP2 (48 kDa), VP4 (29 kDa) and VP3 (33 kDa). During *in vivo* virus maturation pVP2 is processed into VP2 (38 kDa), probably resulting from site-specific cleavage of the pVP2 by a host cell encoded protease (Kibenge et al., 1997). VP2 and VP3 are the two proteins that constitute the single shell of the virion. The B-segment (2.9 kbp) contains one large ORF, encoding the 91 kDa VP1 protein. This protein contains a consensus RNA dependent RNA polymerase motive (Bruenn, 1991). Furthermore, this protein has been reported to be linked to the 5'-ends of the genomic RNA segments (Viral Protein genome-linked, VPg). The nucleotide sequence of internal parts of a large number of IBDV isolates of classical, antigenic variant or very virulent origin has been determined, and deposited in several databases such as GenBank. Furthermore Mundt and Muller (Mundt and Muller, 1995) have determined the 5'- and 3'-termini of several IBDV isolates (CU-1, CU-1M, P-2 and 23/82), and by combining the internal and terminal sequences, Mundt and Muller established

the complete nucleotide sequence of a serotype I A-segment (3261 bp) and B-segment (2827 bp). This provided the way to generate an infectious (recombinant) copy (rIBDV) of IBDV serotype I, by knowing the complete sequence dsRNA sequence of IBDV genome and by using one of several methods to generate infectious copy virus (see for example Boyer et al, Virology 198:415-426, 1994), Mundt and Vakharia indeed produced infectious rIBDV serotype I from cDNA (Mundt and Vakharia, 1996). Full length cDNA of a serotype I IBDV, preceded by a T7 promoter, was thereby used as a template for T7 RNA polymerase using a method described by Weiland and Dreher (Weiland and Dreher, 1989). The *in vitro* generated mRNA, containing a cap-structure at its 5'-end, was subsequently transfected into eukaryotic cells (VERO cells) using a liposome formulation (Lipofectin, GibcoBRL). The supernatant of the transfected cells contained infectious rIBDV after incubation during 36h in the CO<sub>2</sub> incubator at 37° C (Mundt and Vakharia, 1996; (WO 98/09646)). In addition, Lim et al. introduced two amino acid mutations (D279N and A284T) into the cDNA of vvIBDV isolate HK46 (Lim et al., 1999). These mutations were most probably based on data of Yamaguchi et al. (Yamaguchi et al., 1996), which showed that these specific mutations were found in two independent experiments in which very virulent IBDV isolates lost their very virulent character by adaptation and growth on primary CEF cells. Lim et al obtained a rIBDV isolate which possessed the phenotype of a CEF-culture adapted isolate, i.e. a rIBDV isolate which can be propagated, i.e. is able to infect, multiply and be released for further replication, in vvIBDV non-permissive cells such as CEF cells. Note worthy, Lim et al. were unable to produce an infectious vvIBDV isolate using the unmodified cDNA of the HK46 isolate (Lim et al., 1999). Furthermore, although cDNA of IBDV can be used to produce infectious IBDV, the exact mechanism of replication has not been elucidated yet. Data exist which are in support of a semi-conservative genome replication model for Birnaviridae (Bernard, 1980; Mertens et al., 1982).

[0005] Now and then IBDV variants are detected in the field or are created in cell-culture in the laboratory (Muller, 1987) that are genetic re-assortments of serotype I and II strains of IBDV, in that they contain one genomic segment derived from the one serotype, and another segment derived from the other serotype. Such segment reassorted (srIBDV) strains (also called chimeric IBDV) not only occur in nature, but have recently been generated from cDNA as a well, by Vakharia and Mundt (WO 98/09646). Vaccination using attenuated field isolates worked sufficiently well until antigenic derivatives were found in the Delaware region of the USA starting in 1985 (isolates Del A, D, G and E) (Snyder, 1990). These field isolates were missing an important virus neutralizing epitope. The change of this epitope is characterized by the lack of binding of the virus neutralizing monoclonal antibody (Mab) B69 (Snyder et al., 1988a). The antibodies induced by vaccination with classical IBDV vaccines appeared to be less protective against these antigenic IBDV variants. Inactivated vaccines based upon antigenic IBDV variants were subsequently produced and were found to protect effectively against these antigenic variants of IBDV. After the Delaware variant a second antigenic variant IBDV was isolated. This variant was recovered from the Delmarva region (USA) and was referred to as the GLS variant. The GLS variant is characterized by the absence of epitopes for both the virus neutralizing Mab B69 and R63 (Snyder et al., 1988b). After identifying these antigenic variants, a large survey was performed within the USA by using a panel of nine Mabs against IBDV. This survey yielded an additional antigenic variant: the DS326 variant. This antigenic variant is characterized by the absence of epitopes for Mab 179 and BK44, in addition to those for Mabs B69 and R63 (Snyder, 1990). No further reports of antigenic variants have been published in the USA or in other parts of the world. Whether this is due to non-existence of new variant IBDV isolates or whether new antigenic variants just have not been detected due to the lack of extensive surveys or the lack of discriminating monoclonal antibodies is unclear.

[0006] The nucleotide sequence of the polyprotein encoding part of the A-segment of the Del, the GLS and the DS326 antigenic variant IBDV isolates has been determined (Vakharia et al., 1994). Most of the amino acid changes were found in a specific region of the VP2 protein, the so-called hypervariable region. Furthermore it was found that the epitopes which are capable of inducing neutralizing antibodies are conformation dependent and are clustered in the hypervariable region. This region consists of a domain with a high hydrophobicity index (amino acid 224 to 314 of pVP2, corresponding with amino acid 224 to 314 of the polyprotein) which is flanked by two small hydrophilic regions, each spanning about 14 amino acids (Vakharia et al., 1994; Heine et al., 1991). Amino acid substitution both within the hydrophobic region and within the hydrophilic regions might be involved in the antigenic variant character of these isolates.

[0007] After the problems caused by the antigenic variant IBDV isolates in the USA, the poultry industry in Europe was affected by very virulent IBDV (vvIBDV) isolates (Berg et al., 1991; Chettle et al., 1989). The vvIBDV isolates cause more severe clinical signs during an outbreak and are able to break through levels of antibodies which are protective against classical IBDV isolates. The molecular determinants which distinguish vvIBDV from classical IBDV isolates are not exactly known. It is known however, that the pathogenicity of cell culture adapted very virulent IBDV isolates is severely reduced, compared with the non-adapted parental isolates (Yamaguchi et al., 1996). The correlation between CEF-adaptation and loss of the very virulent phenotype is likely to be due to the change in target cell tropism of the adapted virus. This change in cell tropism may be due to the loss of bursa cell receptor binding capability of the cell culture adapted very virulent IBDV isolate. Another possibility is that the cell culture adapted very virulent IBDV isolate is able to infect non-bursa cells, resulting in large reduction of IBDV load in the primary target cells (bursa cells). Form the published results (Yamaguchi et al., 1996), it is clear that a recombinant IBDV (rIBDV) which is based upon

the cDNA of a cell culture adapted very virulent isolate will never yield a vaccine which meets the demands of being able to break through high levels of maternal antibodies and induce a high enough immune response.

[0008] No specific antibodies, that exclusively recognize the vvIBDV isolates have been described yet (Eterradossi et al., 1997)). The lack of discriminating antibodies makes direct diagnosis difficult. Most attention has been given to sequence comparison between the hypervariable region of VP2 of classical isolates and of very virulent isolates. Sequence analysis of the vvIBDV isolate UK661 showed that only three unique (i.e. not found in non-vvIBDV isolates) amino acid substitution are present within the hypervariable region of the VP2 protein. One amino acid substitution is present within the remaining part of the pVP2 protein, while 5 unique amino acid mutations are present within the VP4 encoding part of the polyprotein and 6 in the VP3 encoding part. (Brown and Skinner, 1996). The smaller ORF of the UK661 isolate A-segment, encoding the VP5 protein, contains 2 unique amino acid substitutions. Additionally 16 unique amino acid substitutions were found in the VP1 protein encoded by the B-segment of this vvIBDV isolate. The virulent phenotype of the vvIBDV might be influenced by each of the found amino acid substitutions, and even (silent) nucleotide substitutions within the coding or non-coding parts of either the A- or B-segment may contribute to the altered phenotype of the vvIBDV isolates in comparison with the classical or antigenic variant isolates. Serial passage on embryonated eggs of a vvIBDV isolate (OKYM) resulted in the appearance of a derivative isolate (OKYMT) which is able to grow on Chicken Embryo Fibroblast (CEF) cells and has lost its virulence. This adaptation was reported to be the result of 7 nucleotide substitutions in the polyprotein encoding part of the genome. Whether additional nucleotide substitutions (or deletions) were present in remaining parts of the A- or B-segment (e.g. untranslated regions, VP1 encoding region, and VP5 encoding region) was not determined (Yamaguchi et al., 1996). The reported nucleotide substitutions result in 5 amino acid substitutions. Three of these amino acid substitutions were located in the hydrophobic part of the hypervariable region (I256T, D279N, A284T) of VP2, one in the hydrophilic part located downstream of the hypervariable region (S315F) of VP2, and one in VP3 (A805T) (Yamaguchi et al., 1996). In an independent experiment, Yamaguchi et al. found that the adaptation of vvIBDV isolate TKSM into TKSMT resulted also in the A284T and D279N substitutions. The A284T substitution correlated in their analysis completely with adaptation onto CEF cells and loss of virulence. The D279N substitution was also present in both CEF-adapted vvIBDV isolates (OKYMT and TKSMT) and is potentially also important for growth on CEF cells and loss of virulence. The non-CEF adapted, classical IBDV isolate GBF-1 has on the other hand an asparagine at position 279, in combination with alanine at position 284 and cannot grow on CEF cells, so the single substitution D279N does not account loss of virulence and growth on CEF cells. The amino acid changes in the VP2 apparently allow the modified IBDV to propagate on cells which do not have a receptor for wild type IBDV. Cells possessing a wild-type IBDV receptor such as bursa cells are susceptible for classical and vvIBDV isolates. Recently it was shown that amino acid substitution, A284T in combination with D279N is indeed enough to turn a non-CEF-adapted very virulent IBDV isolate into a CEF-adapted isolate. Lim et al. introduced these two amino acid substitutions into the A-segment cDNA of vvIBDV isolate HK46 (Lim et al., 1999). After transfection of this cDNA, Lim et al obtained a rIBDV isolate which possessed the phenotype of a CEF-culture adapted isolate, i.e. a rIBDV isolate which is able to infect and multiply in CEF cells. The virulence of this rIBDV isolate was not assessed in chickens. Note worthy, Lim et al. were unable to produce a recombinant infectious vvIBDV isolate using the unmodified cDNA of the HK46 isolate (Lim et al., 1999).

[0009] The goal of vaccination against IBD is prevention of subclinical and clinical IBD and the economic aspects of each. Effective vaccination for IBD can be divided into the following categories:

[0010] Protection of the developing bursa in broilers, breeders and layers.

[0011] Prevention of clinical disease in broilers, breeders and layers.

[0012] Priming and boosting of breeders.

[0013] To minimize the immunosuppressive effects of IBDV, the young chick must be protected. Protection of the very young can be achieved through high enough levels of maternal antibodies passed from the breeder hen to her progeny. Vaccination of the very young chick itself may not be successful since onset of protection after vaccination is between three and five days. When a bird, lacking maternal antibodies against IBDV, is exposed to a pathogenic IBDV field strain, damage will occur within 24-48 hours.

[0014] Generally the early vaccinations of the breeders serve as priming. In most situations, this single vaccination is not considered to be adequate. Boosting is the term commonly associated with the administration of a final IBDV vaccination prior to the onset of lay. This is done to increase the circulating antibody in the hen and hence the maternal antibodies in the progeny. Both inactivated (oil emulsion vaccine) and live vaccines (IBDV) have been used for this purpose. The use of a live vaccine in an older bird will result in an increase of antibodies; however, large variations in antibody titers are often seen. These variations result in progeny becoming susceptible to field challenge from as early as a few days after hatching to 21 days after hatching. The use of inactivated IBDV vaccines gives a higher antibody titer as well as a decrease of variation between antibody titers of birds belonging to the same flock. The levels of maternal antibodies necessary to neutralize IBD vary with the invasiveness and pathogenicity of the field strain. In practical terms, if a very virulent IBDV isolate is present, higher maternal antibody levels are desired (see Table 1 for an overview of virulence of field isolates and strength of vaccines). Yet, for effective vaccination, avoiding interference

with maternal antibodies is needed to induce a good immune response. Clinical IBD is typically seen between three and six weeks of age. The immune response of the chick must be stimulated as the passive protection is declining. The timing of the active vaccination may be estimated by the breeder or chick titer and the half-life of antibodies of approximately 3.5 days (De Wit and Van Loon, 1998; Kouwenhoven and Van den Bos, 1995). The levels of maternal antibodies tend to vary within a population. This variation might be a result of variation in the antibodies levels of the breeder hen. Also the mixing of progeny from several breeder flocks (e.g. combination of breeders of different age; breeders vaccinated with live vaccine and those with oil emulsion vaccine) results in variation of IBDV antibodies between chick belonging to the same flock. If the coefficient of variation (CV) in mean maternal antibody titers is too wide, it may be recommended to vaccinate twice (with a 10-day interval) or to vaccinate early with a hot vaccine (in the presence of a high antigenic pressure).

[0015] The average titer of antibody against IBDV in a flock will decline in time (Fig. 1). As a result of the decrease in average antibody titers, an immunity gap will occur. The best results are obtained if the immunity gap is as short as possible and is as early as possible, with a minimum of 2 weeks after hatching. There should be at least sufficient immunity after active vaccination at the age of 4 weeks, since many handlings occur in the houses from that time point with risks of introducing field virus. Therefore farmers like to vaccinate at 2 weeks or even before. Intermediate vaccines are often unable to break through the average IBDV antibody titer of the broiler at two weeks after hatching (Fig. 1). If there is a high variation in mean maternal antibody titers, some chicks will be effectively vaccinated with intermediate vaccines, others not. To circumvent those problems, hot vaccines are being used. A drawback of usage of hot vaccines is that the bursa of chickens with low to moderate maternal antibody titers will be (partly) damaged.

[0016] There is a wide variety of IBDV vaccines available. Important aspects in vaccination strategies are the ability of the virus to replicate in the face of maternal antibody (invasiveness of the vaccine) and the spectrum of antigenic content (including antigenic variants). The ability of a vaccine virus to replicate in the face of maternal antibodies allows live vaccines to be categorized into three main groups: mild, intermediate, and intermediate plus or hot vaccines (see Table 1). The initial vaccines for IBD were derived from classical IBDV isolates. These vaccines were moderately pathogenic IBDV strains with low passage numbers in embryonated eggs. These were often used in breeder programs to induce high levels of circulating antibodies. However, when given to a young bird with moderate or low levels of maternal antibodies, these vaccines could cause extensive bursal atrophy resulting in immunosuppression. Mild vaccines were subsequently developed to be used in these young birds. The attenuation of classical IBDV was done in tissue culture systems. Traditionally, attenuated strains for vaccines are generated by adapting IBDV strains to chicken embryoblast (CEF) cells or other appropriate cells or cell lines through serial passages. These vaccines are not immunosuppressive even when used in birds having no maternal antibodies. However, moderate and high levels of antibodies easily neutralize them. As breeder programs developed (including the use of adjuvant, inactivated vaccines), higher levels of maternal antibodies were generated in progeny. This reduces the effectiveness of these mild vaccines.

[0017] Intermediate strength vaccines were developed to overcome the inadequacies of the mild vaccines. Some of the intermediate vaccines were developed by cloning a field isolate on chicken cell cultures. Intermediate strength vaccines are capable of establishing immunity in birds with moderate levels of maternal antibodies. These vaccines will cause some bursal atrophy in birds without maternal antibodies, but are considered not immunosuppressive.

[0018] Hot (strong) or intermediate plus vaccines were developed after the first outbreaks with vvIBDV. These vvIBDV isolates could break through higher levels of maternal immunity than the vaccines that were on the market at that time. Vaccination with intermediate vaccines came always too late in situations with high infection pressure with vvIBDV. Hot vaccines consist of vvIBDV strains with low to moderate passage in embryonated eggs or bursa derived IBDV of chickens infected with vvIBDV isolates. Adapting vvIBDV on cells traditionally used for the generation of vaccines in general fails, since either these cells are non-permissive for vvIBDV, or, when adapted to said cells, the vvIBDV in question had lost its very virulent character, making it useless for hot or intermediate plus vaccine. Hot or intermediate plus vaccines are desirably able to circumvent maternal immunity at an earlier age than intermediate vaccines but spread more within a flock. If intermediate plus and hot vaccines are used in chickens with moderate to high levels of maternal antibodies, there is no negative side effect on the bursa (Kouwenhoven and Van den Bos, 1995). If these vaccines are used in chickens with low to moderate levels of maternal immunity, this causes depletion of lymphoid cells in the bursa and a severe depletion of peripheral blood-B cells is found (Ducatelle et al., 1995). Although a recovery of bursal function has been observed, these vaccines should be used with precautions.

[0019] Live vaccines must be given in a way in which the virus will preferably reach the bursa where it will quickly multiply and induce an immune response. Possible routes for application of live vaccines include drinking water, spray, subcutaneous and *in ovo*. Inactivated IBD vaccines are used in broiler breeders. They differ in some of the same ways as live vaccines. Their efficacy depends upon the spectrum of antigens they contain. Injectable oil-emulsion products may be given subcutaneously or intramuscularly.

[0020] A continuous monitoring of the field situation using an integrated quality control scheme including serology, can be a valuable tool for continuously adapting preventive vaccination programs to changing epidemiological conditions. Also a continuous follow-up of the epidemiological situation will allow to anticipate the development of major

epidemics (Ducatelle et al., 1995). However, the ability of diagnostic laboratories to monitor IBD with meaningful definitive data is difficult. Serology is important but can be confusing when all birds monitored from commercial broiler flocks have high levels of the same spectrum of circulating antibodies. Field evaluations of broilers to monitor the status of IBD are highly subjective: it is difficult to discriminate antibody titers obtained after vaccination from those induced by IBDV field infections. If it were possible to discriminate between IBDV antibody response to field virus and IBDV vaccination it is possible to have 'early warning' systems and to start IBDV eradication programs if desired. Only when there is a known difference between the antibody response to the used IBDV vaccine and IBDV field isolates, defined conclusion about whether (sub)clinical signs of IBDV are the result of live IBDV vaccination or of IBDV field isolates can be made.

[0021] The invention provides infectious recombinant Infectious Bursal Disease Virus (rIBDV) essentially incapable of growing in a cell that is not derived from a bursa cell or another cell comprising a wild-type IBDV receptor (a non-bursa-cell). A bursa is lymphoid organ, mostly comprising cells that are related to the immune system. In particular, it comprises lymphocytes or lymphocyte precursor cells of sometimes the T-cell- but mainly the B-cell-type, and cells derived thereof, in close relation with monocytes or monocyte derived cells such as macrophages, and also with follicular dendritic cells and antigen presenting cells. In particular, the invention provides rIBDV that is essentially incapable of growing in a cell not listed among above bursa cells or cells derived thereof, such as dendritic cells, monocytes, lymphocytes or cells derived thereof. Herewith the invention provides an rIBDV having retained an important characteristic, in that, in comparison with commonly attenuated IBDV strains, it can not or only little grow in non-bursa cells, such as the well known CEF, QM5 or VERO cells, or other cells that are commonly used for propagating attenuated strains of IBDV. In particular, the invention provides an rIBDV essentially incapable of growing in a non-B-cell derived cell. Essentially incapable of growing herein means that the isolate in question is not or only little capable to infect, multiply or be released for further replication. No such rIBDV isolates exist prior to this invention, all previous rIBDV isolates known grow in non-bursa-cell derived cells such as CEF cells (W098/09646; Lim et al., 1999), thereby for example having lost those very virulent characteristics essential for maintaining in a vaccine strain designed to face above identified problems.

[0022] In a preferred embodiment the invention provides an infectious rIBDV having retained at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus (vvIBDV) needed to provide protection against vvIBDV. In particular, vvIBDV is provided that is essentially incapable of growing in a non-bursa-cell derived cell. In particular, as for example demonstrated in the detailed description, the invention provides an rIBDV essentially incapable of growing in a CEF cell, a VERO cell or a QM5 cell, except of course in those CEF, VERO, QM5, or related cells having been provided with the necessary means (such as transgenic receptor or replication system derived from a bursa-cell) needed for replication of classical or very virulent IBDV.

[0023] Furthermore, the invention provides an rIBDV wherein the amino acid sequence of protein VP2 comprises no asparagine at amino acid position 279, but for example comprises an amino acid particular for a strain with a very virulent character, such as with aspartic acid at amino acid position 279. Such rIBDV strains as provided by the invention have retained at least part of the very virulent characteristics of vvIBDV, as well as an rIBDV according to the invention wherein the amino acid sequence of protein VP2 comprises no threonine at amino acid position 284, but for example comprises an amino acid particular for a strain with a very virulent character, such as with alanine at amino acid position 284.

[0024] In a preferred embodiment, the invention provides an rIBDV according to the invention wherein the amino acid sequence of protein VP2 at least comprises a stretch of amino acids from about position 279 to 289, preferably from about position 229 to 314, most preferably from about position 214 to 328 as found in a vvIBDV isolate such as shown in Table 8.

[0025] The invention furthermore provides a method for obtaining an infectious recombinant copy Infectious Bursal Disease Virus (rIBDV) essentially incapable of growing in a non-bursa-cell derived cell or having at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus (vvIBDV) comprising transfecting at least one first cell with a nucleic acid such as a cDNA or RNA comprising an IBDV genome at least partly derived from a vvIBDV, incubating said first cell in a culture medium, recovering rIBDV from said transfected first cell or said culture medium, propagating said recovered rIBDV in at least one second cell which is permissive for said vvIBDV. A vaccine derived of the recombinant virus as described is also part of this invention. Also a vaccine comprising a chemically or physically inactivated recombinant virus or parts thereof is part of this invention.

[0026] Also the attenuated derivatives of initially produced recombinant very virulent IBDV are part of this invention. Such a virus can be attenuated by known methods including serial passage, removing specific nucleic acid sequences, or by site directed mutagenesis. Physiologically acceptable carriers for vaccines of poultry are known in the art and need not be further described herein. Other additives, such as adjuvants and stabilizers, among others, may also be contained in the vaccine in amounts known in the art. Preferably, adjuvants such as aluminum hydroxide, aluminum phosphate, plant and animal oils, and the like, are administered with the vaccine in amounts sufficient to enhance the immune response to the IBDV. The vaccine of the present invention may also contain various stabilizers. Any suitable

stabilizer can be used including carbohydrates such as sorbitol, mannitol, starch, sucrose, dextrin, or glucose; proteins such as albumin or casein; and buffers such as alkaline metal phosphate and the like. A stabilizer is particularly advantageous when a dry vaccine preparation is prepared by lyophilization. The vaccine can be administered by any suitable known method of inoculating poultry including nasally, ophthalmically, by injection, in drinking water, in the feed, by exposure, and the like. Preferably, the vaccine is administered by mass administration techniques such as *in ovo* vaccination, by placing the vaccine in drinking water or by spraying the animals' environment. When administered by injection, the vaccines are preferably administered parenterally. The vaccine of the present invention is administered to poultry to prevent IBD anytime before or after hatching. Poultry is defined to include but not be limited to chickens, roosters, hens, broilers, roasters, breeders, layers, turkeys and ducks. Examples of pharmaceutically acceptable carriers are diluents and inert pharmaceutical carriers known in the art. Preferably, the carrier or diluent is one compatible with the administration of the vaccine by mass administration techniques. However, the carrier or diluent may also be compatible with other administration methods such as injection, eye drops, nose drops, and the like.

[0027] As explained above, there is need for an IBDV vaccine that can protect against field infections with IBDV, and preferably against very virulent IBDV (vvIBDV). It is clear that vaccines derived from attenuated classical strains and not from very virulent strains will not be able to sufficiently protect. However, as explained above, simply by adapting and cultivating a vvIBDV strain on a cell or cell-line, such as VERO, CEF or QM5, as one skilled in the art would first do in order to obtain a vaccine strain from a vvIBDV strain, reduces its virulent phenotype such that no sufficient protection is to be expected. Therefore, a vaccine strain is needed that has at least partly maintained the very virulent or hot character, in order to provide sufficient protection, however, paradoxically, such a desirable vaccine strain would most likely not be able to be sufficiently or substantially be propagated on appropriate cells, such as non-bursa-cell derived VERO, CEF or QM5, deemed needed to obtain said vaccine. In a preferred embodiment, the invention provides a method wherein said first cell is a non-bursa-cell derived cell non-permissive for said vvIBDV, preferably wherein said first cell has additionally been provided with a helpervirus or a viral protein (herein T7-polymerase is used) derived thereof. With the help of such a cell comprising a properly selected helpervirus, e.g. expressing distinct IBDV or Birna virus viral proteins, or of a cell expressing said IBDV or Birna virus viral proteins, (also called a complementary cell) also now defective or deficient rIBDV can be made.

[0028] The invention therewith also provides a method to generate infectious Infectious Bursal Disease Virus, by combining cDNA sequences derived from very virulent IBDV (vvIBDV) isolates with cDNA sequences derived from either serotype I classical attenuated IBDV isolates, serotype I antigenic variants of IBDV, or serotype II IBDV isolates, wherein said infectious copy recombinant Infectious Bursal Disease Virus having retained at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus has at least retained the incapacity to substantially be propagated on a vvIBDV non-permissive cell such as a QM5 or CEF cell.

[0029] Preferably, a method as provided by the invention provides a vaccine comprising an IBDV genome wherein parts of segments A and/or B derived from a vvIBDV are used combined with parts of segments A and/or B derived from an attenuated IBDV, such as attenuated serotype I or II IBDV. Such a rIBDV is herein also called a mosaic IBDV (mIBDV). Herein we show that (mosaic) vvIBDV can be generated from cDNA by transfection of non-susceptible cells followed by amplification of the cDNA-derived rIBDV on susceptible cells. The method provided herein provides a method to generate vvIBDV from the cloned, full length cDNA of a vvIBDV isolate (see Table 5 and 6). After transfection of QM5 cells with cDNA of vvIBDV it is essential that propagation of the generated vvIBDV virus takes place on cells which are permissive for vvIBDV. These permissive cells can for example be found among Bursa-cell derived cells such as primary bursa cells, in chicken embryo cells, chicken embryo's, or young chickens. Using the method described herein we have for example produced recombinant D6948 (rD6948) using the cDNA derived from the very virulent D6948 IBDV isolate. This rD6948 isolate has the same virulence as the parental D6948 isolate (Table 6).

[0030] Preferably, the invention provides a method wherein a permissive second cell is a primary bursa cell, thereby allowing initial propagation of the desired vaccine virus. As explained above, there is a need for a vaccine capable of breaking through maternal immunity of young chickens at an early stage. A desired vaccine should preferably be able to induce a high level of protection in vaccinated young chickens, and should therefore be as immunogenic as very virulent viruses or be almost as immunogenic.

[0031] The invention furthermore provides a method to engineer recombinant mosaic IBDV (mIBDV) vaccine which has one or more of the desired phenotypes, i.e. i) being able to break through high levels of maternal antibodies in young chickens and being highly immunogenic, ii) having a reduced pathogenicity compared to wild type very virulent IBDV isolates. In one embodiment, the invention provides an infectious mosaic IBDV (mIBDV) comprising a rIBDV wherein at least one genome segment comprises nucleic acid derived from at least two different Birna virus isolates, when preferred wherein at least one of said isolates is a vvIBDV characterised by its incapacity to substantially be propagated on a vvIBDV non-permissive cell such as a VERO, QM5 or CEF cell and/or characterised by its capacity to substantially be propagated on a vvIBDV permissive cell such as a primary bursa cell. For example, the invention provides mIBDV which consists partly of the genome derived from a classical attenuated isolate (such as CEF94) and partly derived from the genome of a vvIBDV isolate (such as D6948). A recombinant mosaic IBDV (mIBDV), made on

the basis of infectious cDNA derived from a very virulent IBDV isolate (D6948), and combined with defined parts of cDNA derived from a cell culture adapted, serotype I, classical IBDV isolate (CEF94) results in a mIBDV isolate which has a reduced pathogenicity compared to wild-type vvIBDV isolates.

[0032] Furthermore specific nucleotide substitutions which either do or do not lead to amino acid mutations, or deletion of specific parts of the IBDV genome again leads to an altered phenotype of the generated mIBDV. For example, the replacement of the pVP2 coding region of CEF-94 cDNA with the corresponding region of cDNA of D6948 yielded plasmid pHB36-vvVP2. This plasmid was subsequently transfected into FPT7 (Britton et al., 1996) infected QM5 cells in combination with pHB-34Z. Supernatant of these transfected QM5 cells was subsequently transferred to fresh QM5 cells. None of these QM5 cells reacted positively in an IPMA using specific antibodies for the VP3 protein of IBDV. On the other hand, primary bursa cells, after being overlaid with supernatant of the transfected cells, reacted positively in the same IPMA. The functional feature of being able to enter permissive cells such as QM5 cells is apparently located in the pVP2 coding region of the A-segment. This invention provides a method to generate recombinant vvIBDV (such as rD6948) having a pVP2 sequence exactly as found in a wild-type vvIBDV (here D6948). All very virulent isolates of which the pVP2 sequences has been described thus far have an alanine at position 284 and cannot or only little be propagated on CEF cells (see Table 7 and 8). On the other hand, when a threonine is present at position 284, propagation on CEF cells is possible, but this is associated with the lack of a very virulent phenotype (see Table 7 and 8). Herein we describe a method to generate infectious recombinant IBDV (rIBDV) having the nucleotide sequence of a wild-type very virulent IBDV isolate, including the alanine codon for amino acid 284, and being unable to be propagated on CEF cells. Furthermore, in our rD6948 isolate we have an aspartic acid present at position 279 in stead of a asparagine commonly found for avirulent IBDV isolates which can be propagated on CEF cells (Table 7 and 8). The rD6948 is truly a very virulent rIBDV, as it is unable to grow on CEF cells (Table 5), and induces similar clinical signs and mortality as wild-type very virulent D6948 isolate (Table 6). Although mIBDV isolate (mCEF94-vvVP2) did, in contrast to the D6948, rD6948 and srIBDV-DACB isolates (also having a functional VP2 protein derived from vvIBDV, see Table 6), not cause any mortality in a 9-days course or body weight loss, it caused the same reduction in bursa weight after 9 days post-infection as the wild-type very virulent D6948 isolate.

[0033] In yet another embodiment, the invention provided a mosaic IBDV according to the invention wherein at least one of said isolates is a serotype II IBDV. Such a mIBDV, preferably lacking at least one immunodominant epitope specific for a serotype I IBDV as well as a (r)D6948 derived vaccine virus such as mD6948-s2VP3C1, also having a functional VP2 protein derived from wIBDV, allowing vaccination with a marker vaccine. Vaccination with a IBDV marker vaccine and subsequent testing with a corresponding diagnostic test enables the discrimination between antibodies induced by the vaccine and by infection with IBDV field isolates. This mIBDV can be differentiated from all other known wild-type IBDV isolates, either belonging to serotype I or serotype II, for example by using a specific set of monoclonal antibodies. The generation of mIBDV from serotype I and II cDNA provides such a mIBDV marker vaccine that induces a serological response in chickens that can be differentiated from the serological response induced by IBDV field strains. The marker vaccine provided by the invention, lacking at least one immunodominant epitope, preferably a serotype I epitope, enables the discrimination between vaccinated and infected animals by means of a diagnostic serologic test. Such a mIBDV marker vaccine is preferably based upon vvIBDV and contains specific sequences originating from classical serotype I or serotype II IBDV. Such a mIBDV marker vaccine has one or more of the following characteristics: i) It induces a protective immune response against vvIBDV field viruses despite high levels of maternal antibodies. ii) It has a reduced pathogenicity compared to vaccines based upon wild-type vvIBDV. iii) It for example misses at least one serotype I specific antigen which enables the serological discrimination of the mIBDV marker vaccine from all serotype I IBDV isolates.

[0034] Also the invention provides a method to produce or generate tailor made vaccines against specific antigenic variants of IBDV by incorporating the specific amino acid changes in a mIBDV vaccine virus. Depending on the composition, these mosaic IBD viruses (mIBDV) possess different phenotypes and different antigenic properties. A specific mutation in one of the viral proteins can have a profound effect on IBDV viability. We found that this is true in case of a single nucleotide substitution, leading to a single amino acid mutation in VP4 (V582A). No rIBDV could be rescued from cDNA when this particular nucleotide substitution was present. Not only mutations within the VP4 encoding region itself, but also mutations or deletions in the region of the cleavage sites (pVP2-VP4 and VP4-VP3) may have a negative effect on replication of rIBDV. Mutations in the other viral proteins, or even deletion of an entire viral protein (i.e. VP5) influences the replication and or virulence as well. Two groups have constructed an VP5 minus rIBDV isolate, by introducing mutations in the cDNA of an CEF-adapted D78 IBDV isolate (Mundt et al., 1997; Yao et al., 1998). Apparently the VP5 protein, which is a non-structural protein, is also a non-essential protein. Yao et al. reported that inactivation of the ORF for VP5 (replacement of the startcodon by a stopcodon) yielded infectious rIBDV (rD78NS which grows to slightly lower titers (*in vitro*) than rD78, while Mundt et al. reported that inactivation of the ORF for VP5 (replacement of the startcodon by a arginine codon) yielded a rIBDV (IBDV/VP5-) which is able to grow to the same titers (*in vitro*) as the parental isolate. Furthermore Yao et al. reported that rD78NS has a decreased cytotoxic and apoptotic effects in cell culture (*in vitro*) and has a delay in replication compared to the parental isolate (*in vivo*), and

failed to induce any pathological lesions or clinical signs of disease in infected chickens.

[0035] Mutations or deletions in the mIBDV cDNA yields a mIBDV with a desired phenotype, i.e. mIBDV which is based on a very virulent isolate but which has a reduced ability to replicate and hence an reduced pathogenicity. The introduction of cDNA sequences from a serotype II, cell culture adapted, IBDV isolate (TY89) into the mosaic virus gives us yet another opportunity to generate marker mIBDV vaccine which can be discriminated from wild-type serotype I IBDV, for example by using specific monoclonal antibodies. Such mIBDV can be used to induce an antibody spectrum, which differs from the spectrum induced by IBDV field isolates. This enables the development of a serologic test to determine whether IBDV antibodies are the result of live mIBDV vaccination or of infection with IBDV field isolates. For example, the mCEF94-s2VP3C virus is recognized by serotype II specific VP3 antibody (Mab T75) while it is also recognized by a serotype I specific VP2 antibody (Mab 1.4). This particular rIBDV is, on the other hand, not recognized by a serotype I specific VP3 antibody (Mab B10). No apparent difference is present between the replication of mCEF94-s2VP3C and rCEF94, indicating that the exchange of the VP3C-terminal part does not lead to major changes in replication ability in QM5 cells. When, on the other hand, the complete VP3 encoding region was exchanged we observed a severe reduction in replication ability of the resulting virus (mCEF94-s2VP3). On the other hand, mCEF94-s2VP3N was not reacting with Mab C3 (VP3, serotype I) while it is fully reacting with Mab B10 (VP3, serotype I) and only partially with Mab T75 (VP3, serotype II). Replication of this mosaic IBDV on CEF cells is reduced compared to rCEF94. From the generated mIBDV, based on cDNA derived from serotype I (CEF94) and serotype II (TY89), it is clear that a serological marker based on VP3 has been identified. The replacement of the cDNA of (part of) VP3 of serotype I for the corresponding part of serotype II, leads to an unique combination of IBDV antigens within one mIBDV isolate. An mIBDV isolate based on this combination of antigens can be used as an IBDV-marker vaccine.

[0036] The introduction of the VP3 C-terminal part of TY89 (Serotype II) into the cDNA of D6948 yielded a mosaic IBDV (mD6948-s2VP3C1) which has a reduced virulence (no mortality, no body weight loss) compared to D6948 or rD6948 (Table 6). This mIBDV, or a comparable isolate which is more or less virulent, is also advantageously used as an IBDV marker vaccine to prevent infections with very virulent IBDV field isolates.

[0037] Furthermore, the invention provides using site-specific mutagenesis techniques to introduce any desired nucleotide mutation within the entire genome of mIBDV. Using this technique allows adapting mIBDV vaccines to future antigenic variations by including any mutation that has been found in antigenic variant IBDV field isolates. Furthermore, it is provided by the invention to exchange part of the genomic sequence of IBDV with the corresponding part of a Birna virus belonging to another genus (e.g. DXV, IPNV, OV, TV). Herewith, the invention provides new mosaic Birna (mBirna) viruses which have new characteristics resulting in new recombinant vaccines for IBDV or other Birna viruses. Also the use of cDNA of other Birna viruses (e.g. DXV, IPNV, OV or TV) leads to new IBDV vaccines. In this approach, one or more of the IBDV immunodominant or neutralizing epitopes are exchanged with the corresponding parts of the protein of another Birna virus.

[0038] Of course, the invention also provides a method for producing an rIBDV according to the invention, said vector comprising heterologous nucleic acid sequences derived from another virus, or (micro)organism, whereby r- or mIBDV serves as a vector. For example a method is provided to generate an infectious copy IBDV which expresses one or more antigens form other pathogens and which can be use to vaccinate against multiple diseases. Such an infectious copy IBDV for example comprises a heterologous cDNA encoding a heterologous protein obtained form a pathogen, for example poultry pathogens. Also a method is provided to generate a conditional lethal IBDV deletion mutant which can be used as self-restricted non-transmissible (carrier) vaccine. Such an IBDV deletion mutant is unable to express one of the IBDV proteins, and is phenotypically complemented by supplying the missing protein by other means.

[0039] The invention is further explained in the detailed description without limiting the invention thereto.

#### Detailed description

#### Material and Methods

#### Viruses and cells

[0040] The IBDV isolate CEF94 is a derivative of PV1 (Petek et al., 1973). After receiving the PV1 isolate in our laboratory in 1973, we have further adapted this isolated by repeated passage (> 25 times) on either primary Chicken Embryo Fibroblast (CEF) cells or Bursa cells. The very virulent IBDV isolate D6948 was originally isolated in 1989 by the Poultry Health Service (Doorn, the Netherlands). It was purified by 5 passages in embryonated eggs and one subsequent passage in SPF leghorn chickens. IBDV Serotype II isolate TY89 (McFerran et al., 1980) was maintained in our laboratory by a limited number of passages on CEF cells. Amplification of CEF-adapted isolates of IBDV (CEF94 and TY89) was performed by growing freshly prepared chicken embryo fibroblast (CEF) cells in a tissue culture flask (75 cm<sup>2</sup>) until near confluence. This cell culture was infected with either CEF94 or TY89 (moi = 0.1) and incubated for 48 h at 37° C in a 5% CO<sub>2</sub> incubator. The supernatant of this culture was centrifuged at 6000 g for 15 min. (pelleting

of debris), transferred to clean tubes and subsequently centrifuged at 33.000 g for 3 h. The virus pellet was resuspended in PBS (1% of the initial culture volume). The very virulent IBDV isolate D6948 was propagated in our laboratory in 21-days-old chickens by inoculation of 200 ELD50 (Egg Lethal Doses) per chicken, nasally and by eye-drop. The bursas of Fabricius were collected from the infected chickens three days post infection, and two volumes of tryptose phosphate buffer was added. This mixture was homogenized in a Sorval Omnimixer (3 \* 10 sec, maximum speed) and subsequently clarified by centrifugation (6000 g, 10 min). The supernatant was transferred to clean tubes and extracted three times with freon and once with chloroform. Virus preparations were stored at -70° C until further use. QM5 cells (Antin and Ordahl, 1991) were received from the laboratory of R. Duncan (Dalhousie University, Halifax, Nova Scotia, Canada) and maintained by using QT35 medium (Fort Dodge), in a CO<sub>2</sub> incubator (37° C).

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#### Isolation of viral dsRNA

[0041] The genomic dsRNA was purified from the IBDV particles by digesting the viral proteins with Proteinase K (Amresco, 1.0 mg/ml) in the presence of 0.5 % SDS during 2 h at 50° C. The viral dsRNA was purified by phenol/ chloroform/isoamylalcohol (25:24:1) extraction (two times) and precipitation with ethanol (2.5V) / NaCl (0.1V, 5M, pH4.8) or with 2 M lithiumchloride (Diaz-Ruiz and Kaper, 1978). The RNA was dissolved in DEPC treated water (10 % of the initial volume) and stored at -20° C until further use. The integrity and purity of the viral RNA was checked on an agarose gel.

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#### Rapid Amplification of cDNA ends.

[0042] The extreme 5'-termini of all genomic RNA strands (the coding and non-coding strands of both the A- and B- segment) of isolate CEF94 were determined. We used 2 ug of genomic dsRNA and 10 pmol of strand- and segment specific primers in a total volume of 12 ul, for each determination. After incubation at 95° C for five min. we transferred this mixture onto ice and added 4 ul of Superscript II first strand syntheses buffer (Gibco/BRL), 2 ul of 100 mM DTT and 2 ul of dNTP's (10 mM each). This mixture was subsequently incubated at 52° C for 2 min, after which 1 ul of reverse transcriptase (Superscript II, Gibco/BRL) was added and incubation at 52° C was continued for one hour. The reverse transcriptase was inactivated by the addition of 1 ul of 0.5 M EDTA. The genomic dsRNA was destroyed by the addition of 2 ul of 6 M NaOH and incubation at 65° C for 30 min. For neutralization, 2 ul of 6 M Acetic acid was added, and cDNA was recovered by using a Qiaex DNA extraction kit (Qiagen) and finally dissolved in 6 ul water. In the anchor ligation reaction we used 2.5 ul of the cDNA preparation, 4 pmol of the anchor, 5 ul T4 ligation buffer and 0.5 ul T4 RNA ligase (New England Biolabs). Incubation was performed at room temperature for 16 h and the reaction was stored at - 20° C. To amplify the single stranded cDNA which was ligated to the anchor, we used a nested primer in combination with the anchor primer. The PCR was performed by using the following conditions: 10 pmol of each specific primer, 10 pmol of the anchor primer, 4.5 or 5.5 mM MgCl<sub>2</sub>, 1\* Taq buffer (Perkin Elmer), 50 uM of each dNTP, 3 units of Taq polymerase (Perkin/Elmer), and 4 ul of the RNA ligation mixture as template, in a total volume of 50 ul. The amplification was performed by 35 cycles through the temperature levels of 92° C (45 sec), 57 or 65°C (45 sec), and 72°C (90 sec). The resulting PCR products were agarose gel purified and digested with EcoRI and XbaI and ligated (T4 DNA Ligase, Pharmacia), in a pUC18 vector which had previously been digested with the same restriction enzymes. The resulting plasmids were amplified in *E. coli* and nucleotide sequence analysis was performed by using the M13F and M13R primers.

#### Generation of full length A- and B-segment single stranded cDNA

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[0043] To produce full length single stranded cDNA of both the A- and B-segments of CEF94 and D6948, we used a primer specific for the 3'-terminus of the coding strand in the reverse transcription reaction for initiation of the cDNA synthesis. As template we used 1 ug of genomic RNA and 2.5 pmol of ANCI (A-segment specific primer, Table 2) or BNCI (B-segment specific primer, Table 2) in a total volume of 10 ul. After incubation at 98° C for two min. we transferred this mixture immediately onto ice and added 10 ul of RT-mix containing 2\* Superscript II first strand syntheses buffer (Gibco/BRL), 20 mM DTT, 2 mM of each dNTP and 100 units of Superscript II (Gibco/BRL). In case of the negative control reaction the addition of Superscript II enzyme was omitted. All tubes were incubated at 50° C for 30 min, after which time 0.5 units of RNase H were added and incubation was continued at 37° C for 15 min. Water (80 ul) was added to each tube, and dsRNA and cDNA was purified by a phenol/chloroform/isoamylalcohol (25:24:1) extraction and precipitated by using a standard ethanol/NaAc precipitation protocol. Obtained pellets were dissolved in 20 ul of water and stored at -20° C.

## Amplification of full length cDNA using a PCR based protocol

[0044] The full length single stranded cDNA of both the A-and B-segment were amplified by using PCR. The primers which hybridize to the 3'-terminus of the non-coding strand of the A-segment (T7AC0, Table 2) and B-segment (T7BC1, Table 2) both have a non-hybridizing 5' extension of 24 nt containing a T7 promoter sequence and an EcoRI site. The primers that hybridize to the 5' terminus of the coding strand of the A-segment (ANCO, Table 2) and B-segment (BNC1, Table 2) match exactly. As template we used 5  $\mu$ l of the above mentioned RT reaction and the PCR was performed in the presence of 1\* Expand High Fidelity buffer, 50  $\mu$ M of each dNTP, 0.2 pmol of each primer, 1.5 units of Expand High Fidelity enzyme, and 2.0 mM MgCl<sub>2</sub> (A-segment) or 4.0 mM MgCl<sub>2</sub> (B-segment). Amplification was performed by cycling 35 times between 94° C (15 sec), 58° C (15 sec) and 72° C (5 min) in case of A-segment amplification (A-program) and cycling for 35 times between 94° C (15 sec), 54° C (15 sec) and 72° C (5 min) in cases of B-segment amplification (B-program), using a Biometra T3 thermocycler. The yield of PCR products was checked on a 1.0% agarose gel.

## 15 Cloning and analysis of the generated PCR fragments

[0045] The full length PCR fragments which were generated three times independently from genomic dsRNA, were isolated from the agarose gel by using a Qiaex gel purification kit (Qiagen) and ligated in the pGEM-Teasy (Promega) vector according to the suppliers instructions. The ligated plasmids were used to transform *E. coli* DH5-alpha cells which were subsequently grown under ampicillin selection (100  $\mu$ g/ml) and in the presence of IPTG (0.8 mg per petri-dish) and Bluo-gal (0.8 mg per petri-dish). Plasmid DNA of white colonies was prepared and analyzed by restriction enzyme digestion and agarose gel separation. The nucleotide sequences of the cloned cDNA's were determined by using a ABI310 automated sequencer and A- and B-segment specific primers. The consensus nucleotide sequences of both segments of CEF94 and of both segments of D6948 were determined (Fig. 2) and the corresponding amino acid sequence of the open reading frames was deduced (Fig. 3). By using the cDNA of two independent clones we restored one amino acid mutation present in the A-segment clone (V542A), resulting in pHB-36W, one amino acid mutation in the A-segment clone of D6948 (P677L), resulting in pHB-60, and one amino acid mutation in the B-segment of D6948 (Q291X), resulting in pHB-55. No amino acid mutations were present in the B-segment cDNA clone of CEF94 (pHB-34Z).

## 30 Introduction of a Hepatitis Delta Virus ribozym

[0046] The Hepatitis Delta Virus ribozym was first introduced into the *E. coli* high copy number plasmid pUC-18 by digesting transcription vector 2.0 (Pattnaik et al., 1992) with restriction enzymes *Xba*I and *Sma*I. The resulting 236 bp fragment, which contains the Hepatitis Delta Virus Ribozym and a T7 RNA polymerase terminator, was ligated in the pUC18 vector which previously was digested with *Xba*I and *Sma*I, yielding pUC-Ribo. Plasmids containing the A- and B-segment of CEF94 and D6948 were used as template in a full length PCR using the above described conditions, and primers specific for either the A-segment (T7AC0 and ANCO) and B-segment (T7BC1 and BNC1). The PCR fragments were agarose gel purified (Qiaex), blunt-ended by using T4 DNA polymerase, and subsequently digested with *Eco*RI. The resulting DNA fragments were directionally cloned into the pUC-Ribo vector which previously had been digested with *Sma*I and *Eco*RI. The resulting plasmids were used as template in an *in vitro* transcription-translation reaction (TnT-T7Quick, Promega). The autoradiogram of SDS-PAGE analyses of the translation products revealed three dominant bands pVP2 (48-49 kDa), VP3 (32-33 kDa), and VP4 (28-29 kDa) when pHB-36A (A-segment of CEF94) or pHB-60 (A-segment of D6948) was used as template. One dominant band (VP1 (91 kDa)) was found when we used plasmid pHB-34Z (B-segment of CEF94) or pHB-55 (B-segment of D6948) as template (data not shown).

## 45 Introduction of a genetic tag

[0047] To distinguish infectious virus generated from cloned cDNA from wild-type virus we introduced a genetic tag in the 3'-UTR of the A-segment of IBDV-A isolate. Two nucleotides of pHB-36A were mutated (C3172T and 3T173A) thereby introducing a unique *Kpn*I restriction site (GGTAAC). These mutations were introduced by the method described by Higuchi (1990). A 383 bp fragment of the resulting PCR fragment was ligated (Rapid ligation kit, Boehringer Mannheim) into the full length A-segment clone (pHB-36A) by using two unique restriction sites (*Bgl*II and *Bpl*). The resulting plasmid pHB-36W was amplified in *E. coli*. The genetic tag was present in this full length CEF94 A-segment clone, as could be concluded from sequence analysis and digestion with restriction enzyme *Kpn*I (data not shown). No difference was observed in the resulting protein pattern when either pHB-36A or pHB-36W was used as template in an *in vitro* transcription/translation reaction (data not shown).

## Construction of mosaic A-segment cDNA

[0048] We constructed plasmids containing mosaic IBDV A-segments which partly consisted of cDNA of one isolate (CEF94) and partly of cDNA of another isolate (D6948). To construct these plasmids we have amplified specific parts of cDNA using appropriate IBDV specific or selective primers. The amplified PCR fragment of cDNA of D6948 was subsequently used to replace the corresponding part in plasmids pHB-36W, using restriction endonucleases and T4 DNA ligase (Rapid DNA Ligation, Boehringer Mannheim).

[0049] For the construction of pHB36-vvVP2 (exchange of pVP2 encoding part, Table 4) we have used IBDV specific to generate the mosaic PCR-VP2D fragment (2256 bp, see Fig. 5a). The internal part of this PCR fragment was used to exchange the corresponding part of pHB-36W, using unique sites for restriction enzymes *Eco*RI and *Sac*II. For the construction of plasmid pHB36-vvVP3 (Table 4) we used IBDV specific primers to generate a mosaic PCR-VP3c fragment (2154 bp, see Fig. 5b). The internal part of this PCR fragment was used to exchange the corresponding part of pHB-36W, using unique sites for the *Eag*I and *Kpn*I (genetic tag site) restriction enzymes.

For the construction of plasmid pHB36-vvVP4 (Table 4) we used IBDV specific primers to generate a mosaic PCR-VP4d fragment (2154 bp, see Fig. 5c). The internal part of this PCR fragment was used to exchange the corresponding part of pHB-36W, using the unique site for restriction enzymes for *Eag*I and *Dra*II.

[0050] plasmids pHB36-vvVP2, -vvVP3, and -vvVP4 were partly analysed by nucleotide sequence determination to conform that no unintended mutations were introduced during the described manipulations.

## 20 Introduction of a serological marker

[0051] To obtain the cDNA of the A-segment of a serotype II IBDV isolate we generated single stranded cDNA of TY89 as described above, by using the ANC1 primer. The coding region of the VP3 protein was subsequently three times independently amplified in a PCR by using 2 ml of RT-material, 1\* Taq buffer, 50 uM of each dNTP, two IBDV serotype II specific primers (0.2 pMol each), 1.5 units of enzyme, and 3.0 mM MgCl<sub>2</sub> in a 0.1 ml reaction volume. Amplification was performed by cycling 35 times between 94° C (15 sec), 52° C (15 sec) and 72° C (1 min). The resulting 956 bp fragment was cloned in the pGEM-TEasy vector and the consensus nucleotide sequence was determined (Fig. 2a). One of the isolated plasmids contained the TY89 VP3 consensus sequence (pSV-VP3-TY89, Fig. 4) and was used as template to generate a 893 bp PCR fragment (see Fig. 5d). This PCR fragment was subsequently used to replace the corresponding part of plasmid pHB36W-vvVP3, by using the artificially introduced *Kpn*I (nt 3175) and *Sac*II (nt 1760) restriction sites in both plasmid pSV-VP3-TY89 and pHB36W-vvVP3. The resulting plasmid (pHB36-vvVP3, see Fig. 5d) encodes the N-terminal 722 amino acids of the CEF94 polyprotein and the 290 C-terminal amino acids of the TY89 polyprotein. The intended exchange was confirmed by nucleotide sequence analysis.

[0052] The same approach was used to exchange the C-terminal half of the coding sequence of the VP3 protein. In stead of the artificially introduced *Sac*II site, we made use of the *Scal* (nt 2799) site which is naturally present both in the TY89 and in the CEF94 cDNA of the A-segment, in combination with the artificially introduced *Kpn*I site (nt 3172). The resulting plasmid (pHB36-s2VP3C, Table 4) encodes a polyprotein consisting of the N-terminal 890 amino acid of the CEF94 polyprotein, in combination with the C-terminal 122 amino acids of the TY89 polyprotein.

[0053] For the construction of plasmid pHB36-s2VP3N (see Table 4) we have replaced the *Scal* (nt 2799) - *Kpn*I (nt 3172) part of plasmid pHB36-s2VP3 with the corresponding part of plasmid pHB-36W. Using the specific restriction endonucleases *Scal* and *Kpn*I, and T4 ligase. The nucleotide sequence of plasmid pHB36-s2VP3N was conformed by sequence analysis.

[0054] For the introduction of the C-terminal encoding part of the VP3 protein of IBDV isolate TY89 into the cDNA of isolate D6948 we have exchanged part of plasmid pHB-60 (nt 1760 -> nt 3260) with the corresponding part of plasmid pHB36-s2VP3C. Plasmid pHB36-s2VP3C was digested with restriction enzymes *Sac*II and *Xba*I and a 1735 bp fragment was recovered from an agarose gel by Qiaex gel extraction kit (Qiaex). This DNA fragment was ligated in the 4440 bp vector fragment of pHB-60 which had previously been digested with the same restriction enzymes. The resulting plasmid (pHB60-s2VP3C1, Table 4) contains cDNA derived from IBDV isolate D6948 (nt 1 to 1760), CEF94 (nt 1760 to 2799 and nt 3175 to 3260), and TY89 (nt 2799 to 3175).

## 50 Transfection of QM5 cells

[0055] QM5 cells, grown to 80% confluence in 60 mm dishes, were infected with Fowl Pox T7 (FPT7) (Britton et al., 1996) one hour prior to transfection. FPT7 infected QM5 cells were subsequently washed once with 5 ml QT-35 medium and incubated with 2 ml fresh Optimem 1 (Gibco/BRL) two times during 15 min. In the mean time, DNA (2.0 to 4.0 ug) was incubated in 0.5 ml Optimem 1 supplemented with 25 ul LipofectAMINE (Gibco/BRL) and kept at room temperature for at least 30 min. The washed QM5 cells were covered with 4 ml of Optimem 1, the DNA/LipofectAMINE transfection mixture was added and the cells were stored for 18h in a 5.0% CO<sub>2</sub> incubator at 37°C.

## Detection of recombinant IBDV after transfection of QM5 cells

[0056] Transfected QM5 cells were washed once with PBS after the transfection. Infectious recombinant IBDV (rIBDV) was recovered from transfected QM5 cells by covering them with 4 ml of QT-35 medium supplemented with 5% fetal calf serum and 2% of an antibiotic mix (1000 U/ml Penicillin, 1000 ug/ml Streptomycin, 20 ug/ml Fungizone, 500 ug/ml Polymixin B, and 10 mg/ml Kanamycin) and incubation for 24 h at 37° C (5.0% CO<sub>2</sub>). The supernatant was filtered through a 200 mM filter (Acrodisc) to remove FPT7 virus and was subsequently stored at -70° C or used directly for quantitation of rIBDV. Recombinant mosaic IBDV (mIBDV) which contains at least the pVP2 from vvIBDV isolate D6948 is unable to re-infect QM5 cells (see Table 5). Therefore, supernatant of transfection experiments which contained D6948 pVP2 encoding cDNA were used to infect 11-days-old, embryonated eggs via the chorioallantoic membrane (CAM) route. To determine the presence of infectious IBDV, the embryo's were collected five days post-infection, homogenized by using a Sorval Omnimixer (3 \* 10 sec, max. speed) and assayed for the presence of IBDV proteins in a IBDV protein specific Elisa.

## 15 Serological differentiation of recombinant mosaic IBDV (mIBDV)

[0057] Different monoclonal antibodies were used to detect recombinant mosaic IBDV (mIBDV) that contained part of the TY89 VP3 or the complete TY89 VP3. The mIBDV's were used to infect QM5 or primary bursa cells and incubated for 24h (QM5 cells) or 48h (primary bursa cells) in a 5% CO<sub>2</sub> incubator at 37° C or 39° C, respectively. The infected cells were subsequently fixed and an immunoperoxidase monolayer assay (IPMA) was performed by using monoclonal antibodies which are either specific for VP2 of serotype I IBDV (Mab 1.4), or specific for VP3 of serotype II (Mab T75), or specific for VP3 of serotype I (Mab B-10 or C-3).

## 25 Virulence of rIBDV in young SPF chickens

[0058] To evaluate the degree of virulence of the generated rIBDV, srIBDV, and mIBDV isolates we have inoculated 12 groups (10 21-days old SPF chickens) with these viruses. Each chicken received nasally and by eye-drop 1000 ELD50 IBDV, with exception with the negative control group, which received only PBS. The animals were monitored for clinical signs and dead chicks were removed each day. At 9 days post infection, all the chicks from the negative control groups and all the surviving chicks from groups in which mortality had occurred, were bled (5 ml) and euthanized for necropsy. From the other groups, 6 chicks were bled (5 ml) and taken for necropsy at day 9 post infection, whereas the remaining 4 were bled (5 ml) and taken for necropsy at day 15 post infection. Bursa and body weight was determined of all chicks which had been euthanized at day 9 post infection

## 35 Results

## Nucleotide sequence determination of the 5'-termini.

[0059] One group has reported the 5'- and 3'-terminal sequences of the segmented dsRNA genome of IBDV (Mundt and Muller, 1995). To verify the terminal sequence of the genome of IBDV and to be able to produce the exact cDNA sequence of a single IBDV isolate we have determined the 5' terminal sequences of both the coding and non-coding strands of the two genomic segments of CEF94, a Chicken Embryo Fibroblast (CEF) adapted, classical isolate of IBDV, by using the RACE (Rapid Amplification of cDNA Ends) technique (Frohman et al., 1988). The RACE analysis was performed in duplicate on each of the four 5'-termini of the CEF94 genome. The resulting sequence data (Table 3) show that the length of the 5'-termini of the coding strands was the same in all cases. Furthermore we found that the nucleotide sequence was identical, except for the last nucleotide which varied in a few clones. This is in contrast to the sequence data of the 5'-termini of the non-coding strands, which varied in length considerably. We also found that the last nucleotide, although preferably a cytosine, varied in some clones similarly to what we found for the 5'-termini of the coding strands. The consensus sequence for the 3'-terminal nucleotide of the A-segment coding strand of CEF94 differs from the nucleotide sequence reported by Mundt and Muller (Mundt and Muller, 1995), i.e. being a cytosine instead of a thymine.

## Generation of plasmids containing full length IBDV cDNA

[0060] Using the sequence data of the 5'-termini we cloned the entire coding and non-coding cDNA sequences of the A-segment and B-segment of classical isolate CEF94 by means of RT-PCR. Using the same procedure and using the same primers we also generated the entire coding and non-coding cDNA of the A- and B-segment of a non-CEF-adapted, very virulent IBDV isolate D6948. The nucleotide sequence of the entire genome of both isolates was deter-

mined three times independently. This sequence information enabled us to generate a consensus nucleotide sequence of both the A- and B-segments of IBDV isolates CEF94 and D6948 (Fig. 2A).

#### 5 Fowlpox T7 polymerase expression system

[0061] One system for generating infectious IBDV virus using *in vitro* synthesized mRNA derived from cDNA of a CEF-adapted IBDV isolate has previously been described (Mundt and Vakharia, 1996). This system is based upon *in vitro* run-off transcription from the T7 promoter which was artificially introduced in front of the cDNA sequences of the A- and B-segments. This RNA is subsequently transfected into VERO cells, after which infectious IBDV virus could be harvested from these cells. One of the drawbacks of this system is that the *in vitro* generated RNA has to contain a 3'-G-*ppp*5'- (cap structure) on its 5'-end in order to get translation of the introduced RNA into the viral proteins, and hence replication of viral RNA. The *in vitro* production of high quality mRNA is both inefficient and expensive as a cap structure has to be present at the 5'-end. Furthermore, expression levels from transfected RNA are generally low due to the short half-life of RNA. To circumvent the drawbacks of generating *in vitro* capped RNA and low expression levels, we have explored the possibility of using an *in vivo* based T7-expression system (Fowlpox T7 polymerase expression system, (Britton et al., 1996) for generation of viral RNA from plasmids containing full length IBDV cDNA.

#### Generating of infectious IBDV using Fowlpox infected cells

[0062] To be able to generate IBDV from cloned cDNA which has the authentic terminal sequences, we introduced the cis-acting Hepatitis Delta Virus (HDV) ribozyme (Chowrira et al., 1994) downstream of the cDNA sequence of the A-and B-segments (Fig. 4). Furthermore we introduced an additional modification in 3' untranslated region of the CEF94 A-segment. By exchanging 2 nucleotides we introduced a unique *Kpn*I endonuclease restriction site in this cDNA. The introduction of this unique restriction site enables us to distinguish between wild-type IBDV and infectious IBDV virus generated from cloned cDNA (genetically tagged rIBDV). As expected, this plasmid yields the same viral proteins in an *in vitro* transcription-translation reaction as the A-segment clone without the genetic tag (data not shown). Plasmid pHB-36W (A-segment CEF94), pHB-60 (A-segment D6948), pHB-34Z (B-segment CEF94), and pHB-55 (B-segment D6948) were used individually to transfet FPT7 infected QM5 cells as described in the Materials and Method section. To analyze whether the transfected QM5 cells expressed IBDV proteins, we performed an IPMA, 24 h after transfection. We used polyclonal antiserum directed either against VP3 (pHB-36W and pHB-60 transfections) or VP1 (pHB-34Z and pHB-55 transfections) in this analysis. About 10 to 50% of the QM5 cell expressed VP3 after transfection with pHB-36W or pHB-60 (data not shown). When B-segment encoding plasmids were used (pHB-34Z or pHB-55) we found that the same percentage of cells (about 10 to 50%) were expressing VP1 (data not shown). Subsequently, we co-transfected combinations of plasmids containing the A- and B-segment cDNA's into FPT7 infected QM5 cells. To screen for infectious recombinant IBDV (rIBDV) in the supernatant of the transfected QM5 cells, we transferred part of the supernatant (10% of the volume) after 18 h onto fresh QM5 cells or onto primary bursa cells. We only could detect rIBDV when A-segments plasmids in combination with B-segments plasmids were used to transfet the QM5 cells. rIBDV could not be detected when supernatant of the cells transfected with A-segment (pHB-60) and B-segment (pHB-55) of D6948 (rD6948) was transferred onto QM5 cells. However, when the co-transfection supernatant of pHB-60 and pHB-55 was transferred onto primary bursa cells or embryonated eggs we were able to show the presence of infectious IBDV (rD6948) in primary bursa cells (after 48h) and in embryonated eggs (after five days). The presence of rIBDV in the first passage was established by using either an IPMA (QM5 cells or primary bursa cells) or an IBDV specific Elisa (embryonated eggs). The generated rCEF94 and rD6948 isolates were amplified in 10-days old embryonated SPF eggs and subsequently used to infect 21-days old SPF chickens (10 chickens per IBDV isolate). The resulting data of the animal experiment (Table 6) shows that the mortality, body weight, bursa weight, and bursa-body weight ratio, caused by rD6948 are the same as the parent very virulent D6948 isolate. Also at necropsy, gross lesions of bursa were as severe for rD6948 as for the parental D6948 isolate (data not shown). From this chicken experiment it is concluded that rD6948 has retained the properties of a very virulent IBDV isolate, and is truly a very virulent rIBDV.

#### 50 Detection of the genetic tag

[0063] Supernatant of rCEF94 infected QM5 cells was harvested and IBDV was isolated by centrifugation as described in the material section. The dsRNA genome was extracted and an A-segment specific primer was used to generate single stranded cDNA, by using reverse transcriptase. The cDNA was subsequently amplified by PCR. The generated PCR fragment was cloned into a high copy number *E. coli* plasmid (pGEM-Teasy, Promega) and was either digested which *Kpn*I or used for nucleotide sequence determination. The presence of the genetic tag in rCEF94 was confirmed in both analyses.

## Identification of a lethal amino acid mutation in VP4

[0064] Plasmid pHB-36 (A-segment CEF94, Table 4) contained a single nucleotide substitution at position 1875 (thymine in stead of a cytosine) compared to the consensus CEF94 A-segment sequence (Fig. 2A). This nucleotide substitution leads to a valine at position 582 of the polyprotein in stead of an alanine, which is encoded by the consensus sequence (V582A, Fig. 3A). As this amino acid mutation is present in the viral protease (VP4), we subsequently checked whether this protease was still able to autocatalytically liberate the viral proteins (pVP2, VP3 en VP4) from the polyprotein. When plasmid-pHB-36 was used as template in a coupled in vitro transcription/translation reaction in the presence of  $^{35}\text{S}$  labeled methionine we found a delayed splicing of the polyprotein (data not shown). Apart from the viral proteins which are found in case of normally spliced polyprotein (pVP2; VP3 and VP4), we found intermediate spliced products (60 kDa: VP4+VP3), and non-spliced polyprotein (data not shown). Although the viral protease (VP4) of clone pHB-36 is able to liberate the structural viral proteins (pVP2 and VP3) from the polyprotein, this clone did not yield rIBDV when using the FPT7 based transfection protocol as described above. Rapid autocatalytic cleavage of the polyprotein is apparently necessary for the generation of infectious rIBDV. We expect that other mutations within VP4 which alter the rate or specificity of the autocatalytic cleavage of the polyprotein will also have a negative effect on viability of the generated rIBDV. Furthermore mutations in the region of the cleavage sites (pVP2-VP4 and VP4-VP3) may also have a negative effect on replication of rIBDV. Any mutation, introduced by modern molecular biological techniques into the cDNA of a very virulent IBDV may enable us to generate rIBDV which has a reduced viability and which can be used as a live or killed IBDV vaccine.

## 20 Generation of segment reassortant IBDV

[0065] Transfection of CEF94 A-segment cDNA (pHB-36W) in combination with D6948 B-segment cDNA (pHB-55) yielded segment reassorted IBDV (srIBDV-CADB) when supernatant of QM5 transfected cells was transferred onto fresh QM5 cells (Table 5). When D6948 A-segment cDNA (pHB-60) was used in combination with CEF94 B-segment cDNA (pHB-34Z) no infectious srIBDV (srIBDV-DACB) could be detected on QM5 cells (Table 5). However, when primary bursa cells were used to assay for the presence of infectious IBDV we found in both cases (srIBDV-CADB and srIBDV-DACB) infected cells after 24h of incubation. Out of the population of primary bursa cells, only lymphoid cells were infected with srIBDV-DACB, while both lymphoid and fibroblast cells were infected in the case of srIBDV-CADB. The srIBDV-DACB isolate induces the same clinical signs as D6948, while the srIBDV-CADB isolate has a comparable virulence as CEF94 (Table 6).

## Construction of mosaic IBDV

[0066] By using modern molecular biological techniques such as those described above, we have created mosaic recombinant IBDV (mIBDV) which exists partly of cDNA derived from CEF94, and partly from D6948 (vvIBDV) or TY89 (a serotype II IBDV isolate). Replacement of the pVP2 protein encoding sequence of CEF94 by the corresponding part of the D6948 yielded only virus (mCEF94-vvVP2) when the supernatant of transfected cells was transferred to cells which are normally susceptible for non CEF-adapted vvIBDV, i.e. primary bursa cells or embryonated eggs. (Table 5). Replacement of the VP3 or VP4 protein encoding sequence of CEF94 with the corresponding part of D6948 yielded mIBDV by using QM5 cells as recipient in the first passage (mCEF-vvVP3 and mCEF-vvVP4 respectively).

[0067] Replacement of the complete VP3 cDNA (290 amino acids) of CEF94 by the corresponding part of the TY89 cDNA yielded a plasmid which encoded a polyprotein consisting of pVP2 and VP4 derived from CEF94 and of VP3 derived from TY89. When this plasmid (pHB36-s2VP3) was used in an in vitro transcription-translation reaction, all the expected proteins, pVP2, VP4 and VP3 were present (data not shown). Transfection of this plasmid in combination with a plasmid (pHB-34Z) containing the B-segment cDNA of CEF94 yielded a mosaic IBDV (mCEF94-s2VP3). Two monoclonal antibodies which are specific for serotype I VP3 (Mab B10 and C3) were unable to recognize this mCEF94-s2VP3, while an antibody which is specific for the serotype II VP3 (Mab T75) did recognize this mosaic virus (Table 5). As expected the mCEF94-s2VP3 was also recognized by a serotype I specific, neutralizing monoclonal antibody directed against VP2 of the CEF94 isolate (Mab 1.4). The TCID<sub>50</sub> on QM5 cells, which was determined 18 hours after transfection, was considerably lower (3 logs) in the case of mCEF94-s2VP3 compared to rCEF94. Furthermore we found that only single QM5 cells were infected by mCEF94-s2VP3 after 24 h. This is in contrast to the plaque forming phenotype of CEF94 and rCEF94 on QM5 cells after 24 h of infection. To generate mIBDV which has the same replication and plaque forming characteristics as rCEF94, but which is still antigenetically different from rCEF94 we subsequently exchanged only the N-terminal part (168 amino acids) or C-terminal part (122 amino acids) of the VP3 of CEF94 by the corresponding sequence of TY89. When these mosaic A-segment plasmid (pHB36-s2VP3N or pHB36-s2VP3C) were transfected in combination with pHB-34Z (CEF94 B-segment) we obtained mosaic IBDVs (mCEF94-s2VP3N or mCEF94-s2VP3C) with replication capabilities in QM5 cells that are equal (mCEF94-s2VP3C)

or slightly reduced (mCEF94-s2VP3N) to those of rCEF94 IBDV (data not shown). Subsequently we checked the recognition of mCEF94-s2VP3N and mCEF94-s2VP3C virus by several Mabs in an IPMA on QM5 infected cells (Table 5). Mab T75 which is specific for VP3 of serotype II also recognized mCEF94-s2VP3C, while the recognition of mCEF94-s2VP3N was slightly reduced. Mab B10, which is specific for VP3 of serotype I did not recognize rCEF94-s2VP3C, while it still recognized mCEF94-s2VP3N. Another Mab (C3) which did not react with mCEF94-s2VP3 infected cells did react with mCEF94-s2VP3C infected cells, although the reaction was reduced compared to QM5 cells infected with rCEF94 (Table 5). mCEF94-s2VP3N was not recognized by Mab C3. The serotype I specific, neutralizing antibody Mab 1.4 which recognizes VP2 recognized, as expected, both mCEF94-s2VP3N and mCEF94-s2VP3C.

[0068] The coding sequence of the C-terminal part of serotype II VP3 (122 amino acids) was also used to replace the corresponding part of the cDNA of D6948. During the exchange we have replaced some D6948 cDNA sequence (encoding for C-terminal part of VP4 and the N-terminal part of VP3, and the 3'-UTR) with the corresponding sequence of CEF-94 (see Fig. 5g). The resulting plasmid (pHB60-s2VP3C1) was, together with pHB-55 (B-segment D6948), transfected into FPT7 infected QM5 cells. Supernatant of these transfected QM5 cells was collected after 24 h and was transferred to embryonated eggs and primary bursa cells. By using monoclonal antibodies we were able to detect infected cells in the monolayer of primary bursa derived cells (see Table 5). mD6948-s2VP3C1 gave the same reaction pattern with the monoclonal antibodies as mCEF-s2VP3C did. Isolate mD6948-s2VP3C1 (1000 ELD50/chicken) was also used to infect 10 SPF chickens (21-days old) to evaluate its virulence. This mIBDV isolate did not cause any mortality in a 9-days course, opposite to the D6948, rD6948 and srIBDV-DACB isolates (Table 6). However, the bursa is severely damaged by this mIBDV, as the bursa-body weight ratio of this group is same as found in the groups which received D6948 or rD6948. This indicates that mD6948-s2VP3C1 is still able to replicate and induce apoptosis in the bursa of Fabricius.

#### LEGENDS TO FIGURES

#### [0069]

- Fig. 1: Antibody titers in broilers having high levels of maternal antibody at day 0.
- Fig. 2a: Nucleotide sequences A-segments
- Fig. 2b: Nucleotide sequences B-segments
- Fig. 3a: Amino acid sequences polyproteins
- Fig. 3b: Amino acid sequence VP1
- Fig. 3c: Amino acid sequence VP5
- Fig. 4: Plasmid drawings
- Fig. 5a: Construction of pHB36-vvVP2
- Fig. 5b: Construction of pHB36-vvVP3
- Fig. 5c: Construction of pHB36-vvVP4
- Fig. 5d: Construction of pHB36-s2VP3
- Fig. 5e: Construction of pHB36-s2VP3C
- Fig. 5f: Construction of pHB36-s2VP3N
- Fig. 5g: Construction of pHB60-s2VP3C1

Table 1:

45	Classification of live IBDV vaccines used to induce active protection in young chickens which are passively protected by maternal IBDV antibodies.		
50	Type of vaccine (live IBD virus)	Ability to induced an immune response when IBDV antibody titers are equal or below	Immunosuppressive
	Mild	50-100	No
	Intermediate	100-200 *	No
55	Strong	500 *	Yes

\* The Animal Health Service (Deventer, The Netherlands) uses an Idexx Elisa value of 128 (2log7) as the maximum titer for the use of live intermediate vaccines and a value of 512 (2log9) for strong vaccines.

Table 2 Primers (oligonucleotides) used for sequence determination, in RT-PCR or PCR reactions. Nucleotides which are unable to hybridize with wild-type IBDV genomes are given in small face. Primers which are specific either for the serotype II (s2) or very virulent (vv) genome are indicated.

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Name	Sequence	Position
Anchor	cacgaattcactatcgattctggatccttc	-
Anchor Primer	gaaggatccagaatcgatag	-
ANC0	GGGGACCCCGCGAACGGATC	A: -1/-18
ANC1	GGGGACCCCGCGAACGG	A: -1/-16
T7AC0	ggaattctaatacgactcaactataGGATAACGATCGGTCTGACCCGG	A: 1/23
BNC1	GGGGGCCCGCAGG	B: -1/-15
T7BC1	ggaattctaatacgactcaactataGGATAACGATGGGTCTGACCCT	B: 1/21

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Table 3 Nucleotide sequence corresponding to the 5'- and 3'-termini of the coding strands of the two genomic segments of IBDV (CEF94). Numbers behind specific sequences indicate the number of times each sequence was obtained.

5'-terminus of the A-segment coding strand	Complementary sequence of the 5'-terminus of the B-segment coding strand	Complementary sequence of the 5'-terminus of the B-segment non-coding strand
5' UGAUACGAUC>>>	>>>CGGGG <sup>3'</sup>	5' UGAUACGAUG>>> (2x)
5' AGAUACGAUC>>>	>>>CGGGGUCCC <sup>3'</sup>	>>>GGAUACGAU>>> (5x)
5' GGAUACGAUC>>>	(7x) >>>CGGGGUCCCU <sup>3'</sup>	>>>GGGGGGCC <sup>3'</sup> (2x)
	>>>CGGGGUCCCCC <sup>3'</sup> (6x)	>>>GGGGGGCCCC <sup>3'</sup> (2x)
	>>>CGGGGUCCCCCU <sup>3'</sup>	>>>GGGGGGCCCCC <sup>3'</sup>
	>>>CGGGGUCCCCCC <sup>3'</sup>	>>>GGGGGGCCCCG <sup>3'</sup>
Consensus 5' GGAUACGAUC>>>	>>>CGGGGUCCCC <sup>3'</sup> (nt 3260) 5' GGAUACGAUG>>>	>>>GGGGGGCCCC <sup>3'</sup> (nt 2827)

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Table 4 Description of the used plasmids

Name	Based on plasmid	Description
pUC18-Ribo	PUC18	Contains the <i>Sma</i> I- <i>Xba</i> I fragment of pTV-2A
pHB-36A	pUC18-Ribo	Contains the consensus cDNA sequence of the A-segment of CEF94 (see Fig. 2a)
pHB-36W	pHB-36A	An artificially introduced <i>Kpn</i> I-site (genetic tag) in the 3'-UTR of the CEF94 A-segment encoding cDNA (Fig. 2)
pHB-36	pHB-36A	Contains a lethal amino acid substitution in the VP4 part of the polyprotein (V582A)
pHB-60	pUC18-Ribo	Contains the consensus cDNA sequence of the D6948 A-segment (see Fig. 2a)
pHB-34Z	pUC18-Ribo	Contains the consensus cDNA sequence of the CEF94 B-segment (see Fig. 2b)
pHB-55	pUC18-Ribo	Contains the consensus cDNA sequence of the D6948 B-segment (see Fig. 2b)
pSV-TY89-VP3	pGEM-Teasy	Contains the consensus cDNA of TY89 encoding the entire VP3 (A-segment, see Fig. 2)
pHB36-vvVP2	pHB-36W	Contains D6948 A-segment cDNA which encodes the entire VP2 (453 amino acids)
pHB36-vvVP3	pHB-36W	Contains D6948 A-segment cDNA which encodes the entire VP3 (289 amino acids)
pHB36-vvVP4	pHB-36W	Contains D6948 A-segment cDNA which encodes the entire VP4 (270 amino acids)
pHB36-s2VP3	pHB-36W	Contains TY89 A-segment cDNA which encodes the entire VP3 (289 amino acids)
pHB36-s2VP3C	pHB-36W	Contains TY89 A-segment cDNA which encodes the C-terminal part (122 amino acids) of VP3
pHB36-s2VP3N	pHB-36W	Contains TY89 A-segment cDNA which encodes the N-terminal part (168 amino acids) of VP3
pHB60-s2VP3C1	pHB-60	Contains cDNA encoding a mosaic polyprotein (D6948 (1-543 AA), CEF94 (544-889 AA), and TY89 (890-1012 AA). The 5'-UTR is derived from D6948, while the 3'-UTR is derived from CEF94. An unique <i>Kpn</i> I-site (genetic tag) is furthermore present in the 3'-UTR

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Table 5 Description of the generated rIBDV, sriBDV, and mIBDV. The ability of these viruses to infect QM5 or primary bursa cells was examined in an immuno peroxidase monolayer assay (IPMA) using either polyclonal serum directed against VP3 or monoclonal antibodies directed against VP2 of IBDV serotype I (1.4), VP3 of serotype II (T-75), or VP3 of serotype I (B10 and C3); nd means not determined.

IBDV virus	Derived from plasmids	B-segment	QM5 cells Bursa	Replication on	Mab's
rCEF94		PHB-34Z	+	+	- + +
rD6948	PHB-36W	PHB-55	+	+	- + +
	PHB-60	PHB-55	-	nd	nd nd
sriBDV-DACB	PHB-60	PHB-34Z	-	nd	nd nd
sriBDV-CADB	PHB-36W	PHB-55	+	nd	nd nd
mCEF94-vvVP2	PHB36-vvVP2	PHB-34Z	-	nd	nd nd
mCEF94-vvVP3	PHB36-vvVP3	PHB-34Z	+	nd	nd nd
mCEF94-vvVP4	PHB36-vvVP4	PHB-34Z	+	nd	nd nd
mCEF94-s2VP3	PHB36-s2VP3	PHB-34Z	+	nd	- + +
mCEF94-s2VP3C	PHB36-s2VP3C	PHB-34Z	+	nd	- + +
mCEF94-s2VP3N	PHB36-s2VP3N	PHB-34Z	+	nd	+ - +
mD6948-s2VP3C1	PHB60-s2VP3C1	PHB-55	-	+	- + +

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**Table 6** Clinical data of 21-day old chickens infected with wild-type, rIBDV, srIBDV or mIBDV isolates (12 groups of 10 chickens). Each chicken was inoculated with 1000 ELD<sub>50</sub> IBDV, except for the negative control group (PBS), and each group was kept in a separate isolator. The bursa and body weight of euthanized chickens was determined at nine days post infection. Standard deviation is given between brackets, together with the number of animals (n) used for determination of the given numbers. The bursa/body weight ratio for each animal was calculated and mean values (standard deviation) per group are given.

IBDV virus	Number of deads (after 9 days)	Body weight (grams)	Bursa weight (grams)	Bursa/Body weight (*1000)
PBS	0	305 (29, n=10)	1.9 (0.4, n=10)	6.1 (1.2)
CEF94	0	341 (16, n=6)	2.0 (0.6, n=6)	6.0 (1.8)
D6948	3	245 (56, n=7)	0.4 (0.1, n=7)	1.7 (0.6)
rCEF94	0	317 (15, n=6)	1.3 (0.5, n=6)	4.2 (1.3)
rD6948	5	261 (24, n=5)	0.4 (0.1, n=5)	1.7 (0.2)
srIBDV-DACB	2	263 (35, n=8)	0.4 (0.1, n=8)	1.5 (0.3)
srIBDV-CADB	0	314 (13, n=6)	1.8 (0.8, n=6)	5.7 (2.7)
mCEF94-vvVP2	0	309 (27, n=6)	0.6 (0.2, n=6)	1.9 (0.4)
mCEF94-vvVP3	0	325 (33, n=6)	2.0 (0.3, n=6)	6.2 (0.7)
mCEF94-vvVP4	0	330 (23, n=6)	1.5 (0.5, n=6)	4.4 (1.3)
mCEF94-s2VP3C	0	320 (11, n=6)	1.3 (0.4, n=6)	4.1 (1.3)
mD6948-s2VP3C1	0	315 (26, n=6)	0.6 (0.2, n=6)	1.9 (0.6)

Table 7: Origin and phenotype of the IBDV isolates

Isolate	Reference	Virulence
D6948	Boot et al., unpublished	Very virulent
rD6948	Boot et al., unpublished	Very virulent
UK661	Brown and Skinner, 1996	Very virulent
5123	Ter Huurne et al., unpublished	Very virulent
96-B4	Ter Huurne et al., unpublished	Avirulent
96-C4	Ter Huurne et al., unpublished	Avirulent
96-C5	Ter Huurne et al., unpublished	Avirulent
96-C6	Ter Huurne et al., unpublished	Very virulent
97-B3	Ter Huurne et al., unpublished	Avirulent
97-B4	Ter Huurne et al., unpublished	Very virulent
97-B5	Ter Huurne et al., unpublished	Very virulent
97-B6	Ter Huurne et al., unpublished	Very virulent
Zoontjes	Ter Huurne et al., unpublished	Very virulent
Hungary	Ter Huurne et al., unpublished	Very virulent
OKYM	Yamaguchi et al., 1996	Very virulent
OKYMT	Yamaguchi et al., 1996	Avirulent
TKSM	Yamaguchi et al., 1996	Very virulent
TKSMT	Yamaguchi et al., 1996	Avirulent
HK46	Lim et al., 1999	Very virulent

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HK46-NT	Lim et al, 1999	Not determined
D6948	<u>214ADDQFESSQXOAGGVTITLPSANIDAI</u> TSLSIGGELVPSRVSQGLLIGDATYLGDTAVITRAVAAADNOLTAGTGTNDLMPNIVIPPTSRITQPTSIKLBETVTS <u>SK9090ADOMWS328</u>	
UK661		
S123		
S6-B4	-P- -L- -P-	V- N- V-
S6-C4		-A-N-
S6-C5		V-
S6-C6		
S7-B3		
S7-B4		
S7-B5		
S7-B6		
Zoontjes		
Hungary		
OKYM		
OKYMT		
TKSM		
TKSMT		
HK46		
HK46-NT		

**Table 8:** Amino acid sequence of the hypervariable region of VP2 of different IBDV isolates. The sequence of the hydrophilic regions (underlined) and the hydrophobic region of very virulent isolate D6948 (amino acid 214 to 328) is used as parental isolate for alignment of the other sequences. Identical amino acid are represented by a dash.

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Annex to the application documents - subsequently filed sequences listing

[0114]

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 35                Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100               105               110  
  
 Gly Asp Gln Met Ser Trp Ser  
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 40                <210> 32  
 <211> 119  
 <212> PRT  
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 45                <221> DOMAIN  
 <222> (1)..(119)  
 <223> /note="Amino acid sequence of the hypervariable  
 region of VP2 of a IBDV isolate"  
  
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 Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Pro Gly Gly Val  
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 Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
 55                20               25               30  
  
 Val Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Val Leu  
 35                35               40               45  
  
 55                Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Thr Val Ile Thr  
 50                55               60  
  
 Arg Ala Val Ala Ala Asn Asn Gly Leu Thr Thr Gly Thr Asp Asn Leu  
 65                70               75               80  
  
 55                Leu Pro Phe Asn Ile Val Ile Pro Thr Asn Glu Ile Thr Gln Pro Ile  
 85                85               90               95

5 Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100 105 110

5 Gly Asp Gln Met Ser Trp Ser  
 115

10 <210> 33

<211> 119

<212> PRT

<213> Infectious bursal disease virus

15 <220>

<221> DOMAIN

<222> (1)..(119)

15 <223> /note="Amino acid sequence of the hypervariable  
 region of VP2 of a IBDV isolate"

20 <400> 33

Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Leu Gly Gly Val  
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20 Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
 20 25 30

25 Ile Gly Gly Glu Leu Val Phe Asn Thr Ser Val Gln Gly Leu Ala Leu  
 35 40 45

25 Asn Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Thr Val Ile Thr  
 50 55 60

30 Arg Ala Val Ala Ser Asp Asn Gly Leu Thr Thr Gly Ile Asp Asn Leu  
 65 70 75 80

35 Met Pro Phe Asn Ile Val Ile Pro Thr Asn Glu Ile Thr Gln Pro Ile  
 85 90 95

35 Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100 105 110

35 Gly Asp Gln Met Ser Trp Ser  
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40 <210> 34

<211> 119

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<213> Infectious bursal disease virus

45 <220>

<221> DOMAIN

<222> (1)..(119)

45 <223> /note="Amino acid sequence of the hypervariable  
 region of VP2 of a IBDV isolate"

50 <400> 34

Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Pro Gly Gly Val  
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50 Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
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55 Val Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Val Leu  
 35 40 45

Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile Thr  
 50 55 60

5 Arg Ala Val Ala Ala Tyr Asn Gly Leu Thr Thr Gly Thr Asp Asn Leu  
 65 70 75 80

10 Leu Pro Phe Asn Ile Val Ile Pro Thr Asn Glu Ile Thr Gln Pro Ile  
 85 90 95

15 Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100 105 110

Gly Asp Gln Met Ser Trp Ser  
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 <213> Infectious bursal disease virus

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<220>  
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 <223> /note="Amino acid sequence of the hypervariable  
 region of VP2 of a IBDV isolate"

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 Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Ala Gly Gly Val  
 1 5 10 15

30

Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
 20 25 30

Ile Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Ile Leu  
 35 40 45

35

Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile Thr  
 50 55 60

Arg Ala Val Ala Ala Asn Asn Gly Leu Thr Ala Gly Thr Asp Asn Leu  
 65 70 75 80

40

Met Pro Phe Asn Ile Val Ile Pro Thr Ser Glu Ile Thr Gln Pro Ile  
 85 90 95

Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100 105 110

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Gly Asp Gln Met Ser Trp Ser  
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 <213> Infectious bursal disease virus

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 Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
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 10 Ile Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Ile Leu  
 35 40 45  
 Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile Thr  
 50 55 60  
 15 Arg Ala Val Ala Ala Asp Asn Gly Leu Thr Ala Gly Thr Asp Asn Leu  
 65 70 75 80  
 Met Pro Phe Asn Ile Val Ile Pro Thr Ser Glu Thr Thr Gln Pro Ile  
 85 90 95  
 20 Thr Ser Ile Lys Leu Val Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100 105 110  
 Gly Asp Gln Met Ser Trp Ser  
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 25 <210> 37  
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 40 Ile Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Thr Leu  
 35 40 45  
 Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile Thr  
 50 55 60  
 45 Arg Ala Val Ala Ala Asn Asn Gly Leu Thr Thr Gly Thr Asp Asn Leu  
 65 70 75 80  
 Met Pro Phe Asn Ile Val Ile Pro Thr Ser Glu Ile Thr Gln Pro Ile  
 85 90 95  
 50 Thr Ser Ile Lys Leu Glu Ile Val Thr Phe Lys Ser Gly Gly Gln Ala  
 100 105 110  
 Gly Asp Gln Met Ser Trp Ser  
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 <223> /note="Amino acid sequence of the hypervariable  
 region of VP2 of a IBDV isolate"  
  
 <400> 38  
 Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Ala Gly Gly Val  
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 Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
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 Ile Gly Gly Glu Leu Val Phe His Thr Ser Val Gln Gly Leu Ile Leu  
 35 40 45  
 Asp Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Thr Thr  
 50 55 60  
 Arg Ala Val Ala Ala Asn Asn Gly Leu Thr Thr Gly Thr Asp Asn Leu  
 65 70 75 80  
 Met Pro Phe Asn Ile Val Ile Pro Thr Ser Glu Ile Thr Gln Pro Ile  
 85 90 95  
 Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
 100 105 110  
 Gly Asp Gln Met Ser Trp Ser  
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 <223> /note="Amino acid sequence of the hypervariable  
 region of VP2 of a IBDV isolate"  
  
 <400> 39  
 Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Ala Gly Gly Val  
 1 5 10 15  
 Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu Ser  
 20 25 30  
 Ile Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Ile Leu  
 35 40 45  
 Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile Thr  
 50 55 60  
 Arg Ala Val Ala Ala Asn Asn Gly Leu Thr Thr Gly Thr Asp Asn Leu  
 65 70 75 80  
 Met Pro Phe Asn Ile Val Ile Pro Thr Ser Glu Ile Thr Gln Pro Ile

85

90

95

Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln Ala  
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 5 Gly Asp Gln Met Ser Trp Ser  
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10 <210> 40  
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 <212> DNA  
 <213> Infectious bursal disease virus

15 <220>  
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 <222> (1)...(3260)  
 <223> /note="cDNA sequence of IBDV A-segment"

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 gatcycagcg atgacraacc tgcaagatca aacccaacag attgttccgt tcatacggag 180  
 cttctgtatc ccaacaaccg gaccggcgtc cattccggac gacaccctrq agaagcacac 240  
 tcttcaggatca gagacctcga cctacaattt gactgtgggg gacacagggt cagggctaat 300  
 tggcttttc cctggwttcc ctggctcaat tgggggtgtct cactacacac tgcagagcaa 360  
 tgggaactac aagttcgatc agatgctct gactgcccag aacctaccgg ccagytacaa 420  
 25 ctactgcagg ctatgtatc ggagtctcact agtgaggatca agcacactyc ctggtggcgt 480  
 ttatgcacta aayggcacca taaacccgt gaccttccaa ggaagcctga gtqaactgac 540  
 agatgttagc tacaatgggt tggatgtctgc aacagccaaac atcaacgaca aaatyyggaa 600  
 cgtccttagta ggggaagggg tmaccgtct cagcttaccc acatcatatg atcttgggt 660  
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 30 atcacagtac caascagggt gggtaacaat cacactgttc tcagcyaaa tygatgccat 840  
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 1320  
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 45 gattgcagga gcattyggct tcaaagacat aatccgggcc mtaaggagga tagctgtgcc  
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 50 agactacctg ctgggcgtat aggcacaggc tgcttcagga actgctcgag cccgcgtcagg  
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 1680  
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 1980  
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 2160  
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 2280  
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 2580  
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 2640  
 25  
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 2700  
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 3120  
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 40  
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 3260

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 <210> 41  
 <211> 3260  
 <212> DNA  
 <213> Infectious bursal disease virus

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 <222> (1)..(3260)  
 <223> /note="cDNA sequence of IBDV A-segment"

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 gatcgacgcg atgacaaccc tgcaagatca aacccaaacag attgtccgt tcatacggag 180  
 ctttctgtatc ccaacaaccc gacccggcgtc cattccggac gacaccctgg agaagc 240

5 tctcaggta gagaccccgaa cctacaattt gactgtgggg gacacagggt cagggctaat 300  
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 tgggaactac aagttcgatc agatgtctt gactgcccag aacctaccgg ccagttacaa 420  
 ctactgcagg ctgtgtatc ggagtctcac agtgaggta agcacacttc ctgggtggcgt 480  
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 agatgttagt tacaatgggt tggatgtctgc aacagccaaac atcaacgaca aaattggaa 600  
 cgtccttagta ggggaaagggg tcaccgtctt cagttaccc acatcatatg atcttgggt 660  
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 10 tgacagcagt gacaggccca gaggcttacac cataactgca gccgatgtt accaattctc 780  
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 aacmarcgag ataaccacgc caatcacatc catcaaactg gagatgtga cctccaaaag  
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 25 cgaggtatgc gcaatctat tccaggtgcc ccagaatccc gtagtcgacg ggattcttgc  
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 1980  
 35 tccactggag actggggagag actacaccgt tggtccaaata gatgtgtct gggacgacag  
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 40 caatgcttgc ggcgagattt agaaagtaag cttttagaaac accaagctcg ccactgcaca  
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 45 ggcaacgttc atcaaaccgtt tccctcacaat tccacgcgac tgggacaggc tcccttaccc  
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 3060  
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 3120  
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25 <210> 42  
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 <212> DNA  
 <213> Infectious bursal disease virus

30 <220>  
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 <222> (1)..(3260)  
 <223> /note="cDNA sequence of IBDV A-segment"

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 40 gatcgacgcg atgacgaaacc tgcaagatca aacccaacag attgttccgt tcatacgag 180  
 ctttctgtatg ccaacaacccg gaccggcgtc cattccggac gacaccctag agaagcacac 240  
 tctcaggatca gagacctcga ctttacaattt gactgtgggg gacacagggt cagggctaat 300  
 tggatgttttc cctgggttcc ctggctcaat tggatgtgttgc cactacacac tgcaagagcaa 360  
 tgggaactac aagttcgatc agatgttccct gactgccccag aaccttacccg ccagctacaa 420  
 ctactgcgg ctgtgtatc ggatgttccac agtggatgtca agcacactcc ctggatgtgt 480  
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 45 agatgttagc tacaatgggt tgatgttcc acacggccaa atcaacgaca aaatcgggaa 600  
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 tgacagcgtt gacaggccca gatgttccac cataactgtca gccgttccat accaatttcc 780  
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 50 gggatgttccat atcttccat tggatgttgc gtttggatgttgc gtaatccca gagctgtggc 960  
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 1020  
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&lt;223&gt; /note="cDNA sequence of IBDV B-segment"

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55  
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	35	40	45
5	Val Gly Asp Thr Gly Ser Gly Leu Ile Val Phe Phe Pro Gly Phe Pro 50 55 60		
	Gly Ser Ile Val Gly Ala His Tyr Thr Leu Gln Ser Asn Gly Asn Tyr 65 70 75 80		
10	Lys Phe Asp Gln Met Leu Leu Thr Ala Gln Asn Leu Pro Ala Ser Tyr 85 90 95		
	Asn Tyr Cys Arg Leu Val Ser Arg Ser Leu Thr Val Arg Ser Ser Thr 100 105 110		
15	Leu Pro Gly Gly Val Tyr Ala Leu Asn Gly Thr Ile Asn Ala Val Thr 115 120 125		
	Phe Gln Gly Ser Leu Ser Glu Leu Thr Asp Val Ser Tyr Asn Gly Leu 130 135 140		
20	Met Ser Ala Thr Ala Asn Ile Asn Asp Lys Ile Gly Asn Val Leu Val 145 150 155 160		
	Gly Glu Gly Val Thr Val Leu Ser Leu Pro Thr Ser Tyr Asp Leu Gly 165 170 175		
25	Tyr Val Arg Leu Gly Asp Pro Ile Pro Ala Ile Gly Leu Asp Pro Lys 180 185 190 195		
	Met Val Ala Thr Cys Asp Ser Ser Asp Arg Pro Arg Val Tyr Thr Ile 195 200 205		
30	Thr Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Xaa Gly Gly 210 215 220		
	Val Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu 225 230 235 240		
35	Ser Xaa Gly Gly Glu Leu Val Phe Gln Thr Ser Val Xaa Gly Leu Xaa 245 250 255		
	Leu Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile 260 265 270		
40	Thr Arg Ala Val Ala Ala Xaa Asn Gly Leu Thr Xaa Gly Thr Asp Asn 275 280 285		
	Leu Xaa Pro Phe Asn Xaa Val Ile Pro Thr Xaa Glu Ile Thr Gln Pro 290 295 300		
45	Ile Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln 305 310 315 320		
	Ala Gly Asp Gln Met Ser Trp Ser Ala Xaa Gly Ser Leu Ala Val Thr 325 330 335		
50	Ile His Gly Gly Asn Tyr Pro Gly Ala Leu Arg Pro Val Thr Leu Val 340 345 350		
	Ala Tyr Glu Arg Val Ala Thr Gly Ser Val Val Thr Val Ala Gly Val 355 360 365		
	Ser Asn Phe Glu Leu Ile Pro Asn Pro Glu Leu Ala Lys Asn Leu Val 370 375 380		
55	Thr Glu Tyr Gly Arg Phe Asp Pro Gly Ala Met Asn Tyr Thr Lys Leu		

385	390	395	400
Ile Leu Ser Glu Arg Asp Arg Leu Gly Ile Lys Thr Val Trp Pro Thr			
405 410 415			
5	Arg Glu Tyr Thr Asp Phe Arg Glu Tyr Phe Met Glu Val Ala Asp Leu	420 425 430	
Asn Ser Pro Leu Lys Ile Ala Gly Ala Phe Gly Phe Lys Asp Ile Ile			
435 440 445			
10	Arg Ala Xaa Arg Arg Ile Ala Val Pro Val Val Ser Thr Leu Phe Pro	450 455 460	
Pro Ala Ala Pro Leu Ala His Ala Ile Gly Glu Gly Val Asp Tyr Leu			
465 470 475 480			
15	Leu Gly Asp Glu Ala Gln Ala Ala Ser Gly Thr Ala Arg Ala Ala Ser	485 490 495	
Gly Lys Ala Arg Ala Ala Ser Gly Arg Ile Arg Gln Leu Thr Leu Ala			
500 505 510			
20	Ala Asp Lys Gly Tyr Glu Val Val Ala Asn Leu Phe Gln Val Pro Gln	515 520 525	
Asn Pro Val Val Asp Gly Ile Leu Ala Ser Pro Gly Xaa Leu Arg Gly			
530 535 540			
25	Ala His Asn Leu Asp Cys Val Leu Arg Glu Gly Ala Thr Leu Phe Pro	545 550 555	560
Val Val Ile Thr Thr Val Glu Asp Ala Met Thr Pro Lys Ala Leu Asn			
565 570 575			
30	Ser Lys Met Phe Ala Val Ile Glu Gly Val Arg Glu Asp Leu Gln Pro	580 585 590	
Pro Ser Gln Arg Gly Ser Phe Ile Arg Thr Leu Ser Gly His Arg Val			
595 600 605			
35	Tyr Gly Tyr Ala Pro Asp Gly Val Leu Pro Leu Glu Thr Gly Arg Asp	610 615 620	
Tyr Thr Val Val Pro Ile Asp Asp Val Trp Asp Asp Ser Ile Met Leu			
625 630 635 640			
40	Ser Lys Asp Pro Ile Pro Pro Ile Val Gly Asn Ser Gly Asn Leu Ala	645 650 655	
Ile Ala Tyr Met Asp Val Phe Arg Pro Lys Val Pro Ile His Val Ala			
660 665 670			
45	Met Thr Gly Ala Leu Asn Ala Xaa Gly Glu Ile Glu Xaa Val Ser Phe	675 680 685	
Arg Ser Thr Lys Leu Ala Thr Ala His Arg Leu Gly Leu Lys Leu Ala			
690 695 700			
50	Gly Pro Gly Ala Phe Asp Val Asn Thr Gly Xaa Asn Trp Ala Thr Phe	705 710 715	720
Ile Lys Arg Phe Pro His Asn Pro Arg Asp Trp Asp Arg Leu Pro Tyr			
725 730 735			
55	Leu Asn Leu Pro Tyr Leu Pro Pro Asn Ala Gly Arg Gln Tyr His Leu		

	740	745	750
5	Ala Met Ala Ala Ser Glu Phe Lys Glu Thr Pro Glu Leu Glu Ser Ala 755 760 765		
	Val Arg Ala Met Glu Ala Ala Asn Val Asp Pro Leu Phe Gln Ser 770 775 780		
10	Ala Leu Ser Val Phe Met Trp Leu Glu Glu Asn Gly Ile Val Thr Asp 785 790 795 800		
	Met Ala Asn Phe Ala Leu Ser Asp Pro Asn Ala His Arg Met Arg Asn 805 810 815		
15	Phe Leu Ala Asn Ala Pro Gln Ala Gly Ser Lys Ser Gln Arg Ala Lys 820 825 830		
	Tyr Gly Thr Ala Gly Tyr Gly Val Glu Ala Arg Gly Pro Thr Pro Glu 835 840 845		
20	Glu Ala Gln Arg Glu Lys Asp Thr Arg Ile Ser Lys Lys Met Glu Thr 850 855 860		
	Met Gly Ile Tyr Phe Ala Thr Pro Glu Trp Val Ala Leu Asn Gly His 865 870 875 880		
25	Arg Gly Pro Ser Pro Gly Gln Leu Lys Tyr Trp Gln Asn Thr Arg Glu 885 890 895		
	Ile Pro Asp Pro Asn Glu Asp Tyr Leu Asp Tyr Val His Ala Glu Lys 900 905 910		
30	Ser Arg Leu Ala Ser Glu Glu Gln Ile Leu Arg Ala Ala Thr Ser Ile 915 920 925		
	Tyr Gly Ala Pro Gly Gln Ala Glu Pro Pro Gln Ala Phe Ile Asp Glu 930 935 940		
35	Val Ala Lys Val Tyr Glu Ile Asn His Gly Arg Gly Pro Asn Gln Glu 945 950 955 960		
	Gln Met Lys Asp Leu Leu Leu Thr Ala Met Glu Met Lys His Arg Asn 965 970 975		
40	Pro Arg Arg Ala Pro Pro Lys Pro Lys Pro Asn Ala Pro Thr 980 985 990		
	Gln Arg Pro Pro Gly Arg Leu Gly Arg Trp Ile Arg Thr Val Ser Asp 995 1000 1005		
45	Glu Asp Leu Glu 1010		
	<210> 48 <211> 1012 <212> PRT 50 <213> Infectious bursal disease virus		
	<220> <221> DOMAIN <222> (1)..(1012) <223> /note="Sequence of IBDV polyprotein CEF94-pp"		
55	<400> 48		

Met Thr Asn Leu Gln Asp Gln Thr Gln Gln Ile Val Pro Phe Ile Arg  
 1 5 10 15

5 Ser Leu Leu Met Pro Thr Thr Gly Pro Ala Ser Ile Pro Asp Asp Thr  
 20 25 30

Leu Glu Lys His Thr Leu Arg Ser Glu Thr Ser Thr Tyr Asn Leu Thr  
 35 40 45

10 Val Gly Asp Thr Gly Ser Gly Leu Ile Val Phe Phe Pro Gly Phe Pro  
 50 55 60

Gly Ser Ile Val Gly Ala His Tyr Thr Leu Gln Ser Asn Gly Asn Tyr  
 65 70 75 80

15 Lys Phe Asp Gln Met Leu Leu Thr Ala Gln Asn Leu Pro Ala Ser Tyr  
 85 90 95

Asn Tyr Cys Arg Leu Val Ser Arg Ser Leu Thr Val Arg Ser Ser Thr  
 100 105 110

20 Leu Pro Gly Gly Val Tyr Ala Leu Asn Gly Thr Ile Asn Ala Val Thr  
 115 120 125

Phe Gln Gly Ser Leu Ser Glu Leu Thr Asp Val Ser Tyr Asn Gly Leu  
 130 135 140

25 Met Ser Ala Thr Ala Asn Ile Asn Asp Lys Ile Gly Asn Val Leu Val  
 145 150 155 160

Gly Glu Gly Val Thr Val Leu Ser Leu Pro Thr Ser Tyr Asp Leu Gly  
 165 170 175

30 Tyr Val Arg Leu Gly Asp Pro Ile Pro Ala Ile Gly Leu Asp Pro Lys  
 180 185 190

Met Val Ala Thr Cys Asp Ser Ser Asp Arg Pro Arg Val Tyr Thr Ile  
 195 200 205

35 Thr Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Pro Gly Gly  
 210 215 220

Val Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu  
 225 230 235 240

40 Ser Val Gly Gly Glu Leu Val Phe Gln Thr Ser Val His Gly Leu Val  
 245 250 255

Leu Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile  
 260 265 270

45 Thr Arg Ala Val Ala Ala Asn Asn Gly Leu Thr Thr Gly Thr Asp Asn  
 275 280 285

Leu Leu Pro Phe Asn Leu Val Ile Pro Thr Asn Glu Ile Thr Gln Pro  
 290 295 300

Ile Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln  
 305 310 315 320

Ala Gly Asp Gln Met Ser Trp Ser Ala Arg Gly Ser Leu Ala Val Thr  
 325 330 335

Ile His Gly Gly Asn Tyr Pro Gly Ala Leu Arg Pro Val Thr Leu Val  
 340 345 350

Ala Tyr Glu Arg Val Ala Thr Gly Ser Val Val Thr Val Ala Gly Val  
 355 360 365  
 Ser Asn Phe Glu Leu Ile Pro Asn Pro Glu Leu Ala Lys Asn Leu Val  
 5 370 375 380  
 Thr Glu Tyr Gly Arg Phe Asp Pro Gly Ala Met Asn Tyr Thr Lys Leu  
 385 390 395 400  
 Ile Leu Ser Glu Arg Asp Arg Leu Gly Ile Lys Thr Val Trp Pro Thr  
 10 405 410 415  
 Arg Glu Tyr Thr Asp Phe Arg Glu Tyr Phe Met Glu Val Ala Asp Leu  
 420 425 430  
 Asn Ser Pro Leu Lys Ile Ala Gly Ala Phe Gly Phe Lys Asp Ile Ile  
 15 435 440 445  
 Arg Ala Ile Arg Arg Ile Ala Val Pro Val Val Ser Thr Leu Phe Pro  
 450 455 460  
 Pro Ala Ala Pro Leu Ala His Ala Ile Gly Glu Gly Val Asp Tyr Leu  
 20 465 470 475 480  
 Leu Gly Asp Glu Ala Gln Ala Ala Ser Gly Thr Ala Arg Ala Ala Ser  
 485 490 495  
 Gly Lys Ala Arg Ala Ala Ser Gly Arg Ile Arg Gln Leu Thr Leu Ala  
 25 500 505 510  
 Ala Asp Lys Gly Tyr Glu Val Val Ala Asn Leu Phe Gln Val Pro Gln  
 515 520 525  
 Asn Pro Val Val Asp Gly Ile Leu Ala Ser Pro Gly Val Leu Arg Gly  
 30 530 535 540  
 Ala His Asn Leu Asp Cys Val Leu Arg Glu Gly Ala Thr Leu Phe Pro  
 545 550 555 560  
 Val Val Ile Thr Thr Val Glu Asp Ala Met Thr Pro Lys Ala Leu Asn  
 35 565 570 575  
 Ser Lys Met Phe Ala Val Ile Glu Gly Val Arg Glu Asp Leu Gln Pro  
 580 585 590  
 Pro Ser Gln Arg Gly Ser Phe Ile Arg Thr Leu Ser Gly His Arg Val  
 40 595 600 605  
 Tyr Gly Tyr Ala Pro Asp Gly Val Leu Pro Leu Glu Thr Gly Arg Asp  
 610 615 620  
 Tyr Thr Val Val Pro Ile Asp Asp Val Trp Asp Asp Ser Ile Met Leu  
 45 625 630 635 640  
 Ser Lys Asp Pro Ile Pro Pro Ile Val Gly Asn Ser Gly Asn Leu Ala  
 645 650 655  
 Ile Ala Tyr Met Asp Val Phe Arg Pro Lys Val Pro Ile His Val Ala  
 50 660 665 670  
 Met Thr Gly Ala Leu Asn Ala Cys Gly Glu Ile Glu Lys Val Ser Phe  
 675 680 685  
 Arg Ser Thr Lys Leu Ala Thr Ala His Arg Leu Gly Leu Lys Leu Ala  
 55 690 695 700

Gly Pro Gly Ala Phe Asp Val Asn Thr Gly Pro Asn Trp Ala Thr Phe  
 705 710 715 720  
 5 Ile Lys Arg Phe Pro His Asn Pro Arg Asp Trp Asp Arg Leu Pro Tyr  
 725 730 735  
 Leu Asn Leu Pro Tyr Leu Pro Pro Asn Ala Gly Arg Gln Tyr His Leu  
 740 745 750  
 10 Ala Met Ala Ala Ser Glu Phe Lys Glu Thr Pro Glu Leu Glu Ser Ala  
 755 760 765  
 Val Arg Ala Met Glu Ala Ala Asn Val Asp Pro Leu Phe Gln Ser  
 770 775 780  
 15 Ala Leu Ser Val Phe Met Trp Leu Glu Glu Asn Gly Ile Val Thr Asp  
 785 790 795 800  
 Met Ala Asn Phe Ala Leu Ser Asp Pro Asn Ala His Arg Met Arg Asn  
 805 810 815  
 20 Phe Leu Ala Asn Ala Pro Gln Ala Gly Ser Lys Ser Gln Arg Ala Lys  
 820 825 830  
 Tyr Gly Thr Ala Gly Tyr Gly Val Glu Ala Arg Gly Pro Thr Pro Glu  
 835 840 845  
 25 Glu Ala Gln Arg Glu Lys Asp Thr Arg Ile Ser Lys Lys Met Glu Thr  
 850 855 860  
 Met Gly Ile Tyr Phe Ala Thr Pro Glu Trp Val Ala Leu Asn Gly His  
 865 870 875 880  
 30 Arg Gly Pro Ser Pro Gly Gln Leu Lys Tyr Trp Gln Asn Thr Arg Glu  
 885 890 895  
 Ile Pro Asp Pro Asn Glu Asp Tyr Leu Asp Tyr Val His Ala Glu Lys  
 900 905 910  
 35 Ser Arg Leu Ala Ser Glu Glu Gln Ile Leu Arg Ala Ala Thr Ser Ile  
 915 920 925  
 Tyr Gly Ala Pro Gly Gln Ala Glu Pro Pro Gln Ala Phe Ile Asp Glu  
 930 935 940  
 40 Val Ala Lys Val Tyr Glu Ile Asn His Gly Arg Gly Pro Asn Gln Glu  
 945 950 955 960  
 Gln Met Lys Asp Leu Leu Leu Thr Ala Met Glu Met Lys His Arg Asn  
 965 970 975  
 45 Pro Arg Arg Ala Pro Pro Lys Pro Lys Pro Lys Pro Asn Ala Pro Thr  
 980 985 990  
 Gln Arg Pro Pro Gly Arg Leu Gly Arg Trp Ile Arg Thr Val Ser Asp  
 995 1000 1005  
 50 Glu Asp Leu Glu  
 1010

<210> 49  
 <211> 1012  
 <212> PRT  
 55 <213> Infectious bursal disease virus

<220>  
 <221> DOMAIN  
 <222> (1)..(1012)  
 <223> /note="Sequence of IBDV polyprotein D6948-PP"  
  
 <400> 49  
 Met Thr Asn Leu Gln Asp Gln Thr Gln Gln Ile Val Pro Phe Ile Arg  
 1 5 10 15  
  
 Ser Leu Leu Met Pro Thr Thr Gly Pro Ala Ser Ile Pro Asp Asp Thr  
 20 25 30  
  
 Leu Glu Lys His Thr Leu Arg Ser Glu Thr Ser Thr Tyr Asn Leu Thr  
 35 40 45  
  
 Val Gly Asp Thr Gly Ser Gly Leu Ile Val Phe Phe Pro Gly Phe Pro  
 50 55 60  
  
 Gly Ser Ile Val Gly Ala His Tyr Thr Leu Gln Ser Asn Gly Asn Tyr  
 65 70 75 80  
  
 Lys Phe Asp Gln Met Leu Leu Thr Ala Gln Asn Leu Pro Ala Ser Tyr  
 85 90 95  
  
 Asn Tyr Cys Arg Leu Val Ser Arg Ser Leu Thr Val Arg Ser Ser Thr  
 100 105 110  
  
 Leu Pro Gly Gly Val Tyr Ala Leu Asn Gly Thr Ile Asn Ala Val Thr  
 115 120 125  
  
 Phe Gln Gly Ser Leu Ser Glu Leu Thr Asp Val Ser Tyr Asn Gly Leu  
 130 135 140  
  
 Met Ser Ala Thr Ala Asn Ile Asn Asp Lys Ile Gly Asn Val Leu Val  
 145 150 155 160  
  
 Gly Glu Gly Val Thr Val Leu Ser Leu Pro Thr Ser Tyr Asp Leu Gly  
 165 170 175  
  
 Tyr Val Arg Leu Gly Asp Pro Ile Pro Ala Ile Gly Leu Asp Pro Lys  
 180 185 190  
  
 Met Val Ala Thr Cys Asp Ser Ser Asp Arg Pro Arg Val Tyr Thr Ile  
 195 200 205  
  
 Thr Ala Ala Asp Asp Tyr Gln Phe Ser Ser Gln Tyr Gln Ala Gly Gly  
 210 215 220  
  
 Val Thr Ile Thr Leu Phe Ser Ala Asn Ile Asp Ala Ile Thr Ser Leu  
 225 230 235 240  
  
 Ser Ile Gly Gly Glu Leu Val Phe Gln Thr Ser Val Gln Gly Leu Ile  
 245 250 255  
  
 Leu Gly Ala Thr Ile Tyr Leu Ile Gly Phe Asp Gly Thr Ala Val Ile  
 260 265 270  
  
 Thr Arg Ala Val Ala Ala Asp Asn Gly Leu Thr Ala Gly Thr Asp Asn  
 275 280 285  
  
 Leu Met Pro Phe Asn Ile Val Ile Pro Thr Ser Glu Ile Thr Gln Pro  
 290 295 300  
  
 Ile Thr Ser Ile Lys Leu Glu Ile Val Thr Ser Lys Ser Gly Gly Gln  
 305 310 315 320

Ala Gly Asp Gln Met Ser Trp Ser Ala Ser Gly Ser Leu Ala Val Thr  
 325 330 335  
 5 Ile His Gly Gly Asn Tyr Pro Gly Ala Leu Arg Pro Val Thr Leu Val  
 340 345 350  
 Ala Tyr Glu Arg Val Ala Thr Gly Ser Val Val Thr Val Ala Gly Val  
 355 360 365  
 10 Ser Asn Phe Glu Leu Ile Pro Asn Pro Glu Leu Ala Lys Asn Leu Val  
 370 375 380  
 Thr Glu Tyr Gly Arg Phe Asp Pro Gly Ala Met Asn Tyr Thr Lys Leu  
 385 390 395 400  
 15 Ile Leu Ser Glu Arg Asp Arg Leu Gly Ile Lys Thr Val Trp Pro Thr  
 405 410 415  
 Arg Glu Tyr Thr Asp Phe Arg Glu Tyr Phe Met Glu Val Ala Asp Leu  
 420 425 430  
 20 Asn Ser Pro Leu Lys Ile Ala Gly Ala Phe Gly Phe Lys Asp Ile Ile  
 435 440 445  
 Arg Ala Leu Arg Arg Ile Ala Val Pro Val Val Ser Thr Leu Phe Pro  
 450 455 460  
 25 Pro Ala Ala Pro Leu Ala His Ala Ile Gly Glu Gly Val Asp Tyr Leu  
 465 470 475 480  
 Leu Gly Asp Glu Ala Gln Ala Ala Ser Gly Thr Ala Arg Ala Ala Ser  
 485 490 495  
 30 Gly Lys Ala Arg Ala Ala Ser Gly Arg Ile Arg Gln Leu Thr Leu Ala  
 500 505 510  
 Ala Asp Lys Gly Tyr Glu Val Val Ala Asn Leu Phe Gln Val Pro Gln  
 515 520 525  
 35 Asn Pro Val Val Asp Gly Ile Leu Ala Ser Pro Gly Ile Leu Arg Gly  
 530 535 540  
 Ala His Asn Leu Asp Cys Val Leu Arg Glu Gly Ala Thr Leu Phe Pro  
 545 550 555 560  
 40 Val Val Ile Thr Thr Val Glu Asp Ala Met Thr Pro Lys Ala Leu Asn  
 565 570 575  
 Ser Lys Met Phe Ala Val Ile Glu Gly Val Arg Glu Asp Leu Gln Pro  
 580 585 590  
 45 Pro Ser Gln Arg Gly Ser Phe Ile Arg Thr Leu Ser Gly His Arg Val  
 595 600 605  
 Tyr Gly Tyr Ala Pro Asp Gly Val Leu Pro Leu Glu Thr Gly Arg Asp  
 610 615 620  
 50 Tyr Thr Val Val Pro Ile Asp Asp Val Trp Asp Asp Ser Ile Met Leu  
 625 630 635 640  
 Ser Lys Asp Pro Ile Pro Pro Ile Val Gly Asn Ser Gly Asn Leu Ala  
 645 650 655  
 55 Ile Ala Tyr Met Asp Val Phe Arg Pro Lys Val Pro Ile His Val Ala  
 660 665 670

Met Thr Gly Ala Leu Asn Ala Tyr Gly Glu Ile Glu Asn Val Ser Phe  
 675 680 685  
 5 Arg Ser Thr Lys Leu Ala Thr Ala His Arg Leu Gly Leu Lys Leu Ala  
 690 695 700  
 Gly Pro Gly Ala Phe Asp Val Asn Thr Gly Ser Asn Trp Ala Thr Phe  
 705 710 715 720  
 10 Ile Lys Arg Phe Pro His Asn Pro Arg Asp Trp Asp Arg Leu Pro Tyr  
 725 730 735  
 Leu Asn Leu Pro Tyr Leu Pro Pro Asn Ala Gly Arg Gln Tyr Asp Leu  
 740 745 750  
 15 Ala Met Ala Ala Ser Glu Phe Lys Glu Thr Pro Glu Leu Glu Ser Ala  
 755 760 765  
 Val Arg Ala Met Glu Ala Ala Asn Val Asp Pro Leu Phe Gln Ser  
 770 775 780  
 20 Ala Leu Ser Val Phe Met Trp Leu Glu Glu Asn Gly Ile Val Thr Asp  
 785 790 795 800  
 Met Ala Asn Phe Ala Leu Ser Asp Pro Asn Ala His Arg Met Arg Asn  
 805 810 815  
 25 Phe Leu Ala Asn Ala Pro Gln Ala Gly Ser Lys Ser Gln Arg Ala Lys  
 820 825 830  
 Tyr Gly Thr Ala Gly Tyr Gly Val Glu Ala Arg Gly Pro Thr Pro Glu  
 835 840 845  
 30 Glu Ala Gln Arg Glu Lys Asp Thr Arg Ile Ser Lys Lys Met Glu Thr  
 850 855 860  
 Met Gly Ile Tyr Phe Ala Thr Pro Glu Trp Val Ala Leu Asn Gly His  
 865 870 875 880  
 35 Arg Gly Pro Ser Pro Gly Gln Leu Lys Tyr Trp Gln Asn Thr Arg Glu  
 885 890 895  
 Ile Pro Asp Pro Asn Glu Asp Tyr Leu Asp Tyr Val His Ala Glu Lys  
 900 905 910  
 40 Ser Arg Leu Ala Ser Glu Glu Gln Ile Leu Arg Ala Ala Thr Ser Ile  
 915 920 925  
 Tyr Gly Ala Pro Gly Gln Ala Glu Pro Pro Gln Ala Phe Ile Asp Glu  
 930 935 940  
 45 Val Ala Lys Val Tyr Glu Ile Asn His Gly Arg Gly Pro Asn Gln Glu  
 945 950 955 960  
 Gln Met Lys Asp Leu Leu Thr Ala Met Glu Met Lys His Arg Asn  
 965 970 975  
 50 Pro Arg Arg Ala Pro Pro Lys Pro Lys Pro Lys Pro Asn Ala Pro Thr  
 980 985 990  
 Gln Arg Pro Pro Gly Arg Leu Gly Arg Trp Ile Arg Ala Val Ser Asp  
 995 1000 1005  
 55 Glu Asp Leu Glu  
 1010

5 <210> 50  
 <211> 290  
 <212> PRT  
 <213> Infectious bursal disease virus  
 10 <220>  
 <221> DOMAIN  
 <222> (1)..(290)  
 <223> /note="Sequence of IBDV polyprotein TY89-PP"  
 15 <400> 50  
 Arg Phe Pro His Asn Pro Arg Asp Trp Asp Arg Leu Pro Tyr Leu Asn  
 1 5 10 15  
 Leu Pro Tyr Leu Pro Pro Thr Ala Gly Arg Gln Phe His Leu Ala Leu  
 20 25 30  
 15 Ala Ala Ser Glu Phe Lys Glu Thr Pro Glu Leu Glu Asp Ala Val Arg  
 35 40 45  
 Ala Met Asp Ala Ala Ala Asn Val Asp Pro Leu Phe Arg Ser Ala Leu  
 20 50 55 60  
 Gln Val Phe Met Trp Leu Glu Glu Asn Gly Ile Val Thr Asp Met Ala  
 65 70 75 80  
 25 Asn Phe Ala Leu Ser Asp Pro Asn Ala His Arg Met Lys Asn Phe Leu  
 85 90 95  
 Ala Asn Ala Pro Gln Ala Gly Ser Lys Ser Gln Arg Ala Lys Tyr Gly  
 100 105 110  
 30 Thr Ala Gly Tyr Gly Val Glu Ala Arg Gly Pro Thr Pro Glu Glu Ala  
 115 120 125  
 Gln Arg Glu Lys Asp Thr Arg Ile Ser Lys Lys Met Glu Thr Met Gly  
 130 135 140  
 35 Ile Tyr Phe Ala Thr Pro Glu Trp Val Ala Leu Asn Gly His Arg Gly  
 145 150 155 160  
 Pro Ser Pro Gly Gln Leu Lys Tyr Trp Gln Asn Thr Arg Glu Ile Pro  
 165 170 175  
 40 Glu Pro Asn Glu Asp Tyr Pro Asp Tyr Val His Ala Glu Lys Ser Arg  
 180 185 190  
 Leu Ala Ser Glu Glu Gln Val Leu Arg Ala Ala Thr Ser Ile Tyr Gly  
 195 200 205  
 45 Ala Pro Gly Gln Ala Glu Pro Pro Gln Ala Phe Ile Asp Glu Val Ala  
 210 215 220  
 Arg Val Tyr Glu Ile Asn His Gly Arg Gly Pro Asn Gln Glu Gln Met  
 225 230 235 240  
 50 Lys Asp Leu Leu Leu Thr Ala Met Glu Met Lys His Arg Asn Pro Arg  
 245 250 255  
 Arg Ala Pro Pro Lys Pro Lys Pro Lys Pro Asn Ala Pro Ser Gln Arg  
 260 265 270  
 55 Pro Pro Gly Arg Leu Gly Arg Trp Ile Arg Thr Val Ser Asp Glu Asp

275

280

285

Leu Glu  
290

5

<210> 51  
 <211> 881  
 <212> PRT  
 10 <213> Infectious bursal disease virus  
  
 <220>  
 <221> DOMAIN  
 <222> (1)..(881)  
 15 <223> /note="Consensus sequence of IBDV VP1, whereby X stands for  
 any amino acid"  
  
 <400> 51  
 Met Ser Asp Xaa Phe Asn Ser Pro Gln Ala Arg Ser Xaa Ile Ser Ala  
 1 5 10 15  
  
 20 Ala Phe Gly Ile Lys Pro Thr Ala Gly Gln Asp Val Glu Glu Leu Leu  
 20 25 30  
  
 25 Ile Pro Lys Val Trp Val Pro Pro Glu Asp Pro Leu Ala Ser Pro Ser  
 35 40 45  
  
 30 Arg Leu Ala Lys Phe Leu Arg Glu Asn Gly Tyr Lys Xaa Leu Gln Pro  
 50 55 60  
  
 35 Arg Ser Leu Pro Glu Asn Glu Glu Tyr Glu Thr Asp Gln Ile Leu Pro  
 65 70 75 80  
  
 40 Asp Leu Ala Trp Met Arg Gln Ile Glu Gly Ala Val Leu Lys Pro Thr  
 85 90 95  
  
 45 Leu Ser Leu Pro Ile Gly Asp Gln Glu Tyr Phe Pro Lys Tyr Tyr Pro  
 100 105 110  
  
 50 Thr His Arg Pro Ser Lys Glu Lys Pro Asn Ala Tyr Pro Pro Asp Ile  
 115 120 125  
  
 55 Ala Leu Leu Lys Gln Met Ile Tyr Leu Phe Leu Gln Val Pro Glu Ala  
 130 135 140  
  
 60 Xaa Xaa Xaa Leu Lys Asp Glu Val Thr Leu Leu Thr Gln Asn Ile Arg  
 145 150 155 160  
  
 65 Asp Lys Ala Tyr Gly Ser Gly Thr Tyr Met Gly Gln Ala Thr Arg Leu  
 165 170 175  
  
 70 Val Ala Met Lys Glu Val Ala Thr Gly Arg Asn Pro Asn Lys Asp Pro  
 180 185 190  
  
 75 Leu Lys Leu Gly Tyr Thr Phe Glu Ser Ile Ala Gln Leu Leu Asp Ile  
 195 200 205  
  
 80 Thr Leu Pro Val Gly Pro Pro Gly Glu Asp Asp Lys Pro Trp Val Pro  
 210 215 220  
  
 85 Leu Thr Arg Val Pro Ser Arg Met Leu Val Leu Thr Gly Asp Val Asp  
 225 230 235 240  
  
 90 Gly Xaa Phe Glu Val Glu Asp Tyr Leu Pro Lys Ile Asn Leu Lys Ser  
 245 250 255

55



Lys Ser Lys Val Gly Ile Glu Gln Ala Tyr Lys Val Val Arg Tyr Glu  
 610 615 620

5 Ala Leu Arg Leu Val Gly Gly Trp Asn Tyr Pro Leu Leu Asn Lys Ala  
 625 630 635 640

Cys Lys Asn Asn Ala Xaa Ala Ala Arg Arg His Leu Glu Ala Lys Gly  
 645 650 655

10 Phe Pro Leu Asp Glu Phe Leu Ala Glu Trp Ser Glu Leu Ser Glu Phe  
 660 665 670

Gly Glu Ala Phe Glu Gly Phe Asn Ile Lys Leu Thr Val Thr Xaa Glu  
 675 680 685

15 Ser Leu Ala Glu Leu Asn Xaa Pro Val Pro Pro Lys Pro Pro Asn Val  
 690 695 700

Asn Arg Pro Val Asn Thr Gly Gly Leu Lys Ala Val Ser Asn Ala Leu  
 705 710 715 720

20 Lys Thr Gly Arg Tyr Arg Asn Glu Ala Gly Leu Ser Gly Leu Val Leu  
 725 730 735

Leu Ala Thr Ala Arg Ser Arg Leu Gln Asp Ala Val Lys Ala Lys Ala  
 740 745 750

25 Glu Ala Glu Lys Leu His Lys Ser Lys Pro Asp Asp Pro Asp Ala Asp  
 755 760 765

Trp Phe Glu Arg Ser Glu Thr Leu Ser Asp Leu Leu Glu Lys Ala Asp  
 770 775 780

30 Ile Ala Ser Lys Val Ala His Ser Ala Leu Val Glu Thr Ser Asp Ala  
 785 790 795 800

Leu Glu Ala Val Gln Ser Thr Ser Val Tyr Thr Pro Lys Tyr Pro Glu  
 805 810 815

35 Val Lys Asn Pro Gln Thr Ala Ser Asn Pro Val Val Gly Leu His Leu  
 820 825 830

Pro Ala Lys Arg Ala Thr Gly Val Gln Ala Ala Leu Leu Gly Ala Gly  
 835 840 845

40 Thr Ser Arg Pro Met Gly Met Glu Ala Pro Thr Arg Ser Lys Asn Ala  
 850 855 860

Val Lys Met Ala Lys Arg Arg Gln Arg Gln Lys Glu Ser Arg Gln Xaa  
 865 870 875 880

45 Xaa

50 <210> 52  
 <211> 881  
 <212> PRT  
 <213> Infectious bursal disease virus

55 <220>  
 <221> DOMAIN  
 <222> (1)...(881)  
 <223> /note="Sequence of IBDV CEF94-VP1"

5 <400> 52  
 Met Ser Asp Ile Phe Asn Ser Pro Gln Ala Arg Ser Thr Ile Ser Ala  
 1 5 10 15  
 Ala Phe Gly Ile Lys Pro Thr Ala Gly Gln Asp Val Glu Glu Leu Leu  
 20 25 30  
 10 Ile Pro Lys Val Trp Val Pro Pro Glu Asp Pro Leu Ala Ser Pro Ser  
 35 40 45  
 Arg Leu Ala Lys Phe Leu Arg Glu Asn Gly Tyr Lys Val Leu Gln Pro  
 50 55 60  
 15 Arg Ser Leu Pro Glu Asn Glu Glu Tyr Glu Thr Asp Gln Ile Leu Pro  
 65 70 75 80  
 Asp Leu Ala Trp Met Arg Gln Ile Glu Gly Ala Val Leu Lys Pro Thr  
 85 90 95  
 20 Leu Ser Leu Pro Ile Gly Asp Gln Glu Tyr Phe Pro Lys Tyr Tyr Pro  
 100 105 110  
 Thr His Arg Pro Ser Lys Glu Lys Pro Asn Ala Tyr Pro Pro Asp Ile  
 115 120 125  
 25 Ala Leu Leu Lys Gln Met Ile Tyr Leu Phe Leu Gln Val Pro Glu Ala  
 130 135 140  
 Asn Glu Gly Leu Lys Asp Glu Val Thr Leu Thr Gln Asn Ile Arg  
 145 150 155 160  
 30 Asp Lys Ala Tyr Gly Ser Gly Thr Tyr Met Gly Gln Ala Thr Arg Leu  
 165 170 175  
 Val Ala Met Lys Glu Val Ala Thr Gly Arg Asn Pro Asn Lys Asp Pro  
 180 185 190  
 35 Leu Lys Leu Gly Tyr Thr Phe Glu Ser Ile Ala Gln Leu Leu Asp Ile  
 195 200 205  
 Thr Leu Pro Val Gly Pro Pro Gly Glu Asp Asp Lys Pro Trp Val Pro  
 210 215 220  
 40 Leu Thr Arg Val Pro Ser Arg Met Leu Val Leu Thr Gly Asp Val Asp  
 225 230 235 240  
 Gly Asp Phe Glu Val Glu Asp Tyr Leu Pro Lys Ile Asn Leu Lys Ser  
 245 250 255  
 45 Ser Ser Gly Leu Pro Tyr Val Gly Arg Thr Lys Gly Glu Thr Ile Gly  
 260 265 270  
 Glu Met Ile Ala Ile Ser Asn Gln Phe Leu Arg Glu Leu Ser Thr Leu  
 275 280 285  
 50 Leu Lys Gln Gly Ala Gly Thr Lys Gly Ser Asn Lys Lys Lys Leu Leu  
 290 295 300  
 Ser Met Leu Ser Asp Tyr Trp Tyr Leu Ser Cys Gly Leu Leu Phe Pro  
 305 310 315 320  
 Lys Ala Glu Arg Tyr Asp Lys Ser Thr Trp Leu Thr Lys Thr Arg Asn  
 325 330 335  
 55 Ile Trp Ser Ala Pro Ser Pro Thr His Leu Met Ile Ser Met Ile Thr

	340	345	350
5	Trp Pro Val Met Ser Asn Ser Pro Asn Asn Val Leu Asn Ile Glu Gly 355 360 365		
	Cys Pro Ser Leu Tyr Lys Phe Asn Pro Phe Arg Gly Gly Leu Asn Arg 370 375 380		
10	Ile Val Glu Trp Ile Leu Ala Pro Glu Glu Pro Lys Ala Leu Val Tyr 385 390 395 400		
	Ala Asp Asn Ile Tyr Ile Val His Ser Asn Thr Trp Tyr Ser Ile Asp 405 410 415		
15	Leu Glu Lys Gly Glu Ala Asn Cys Thr Arg Gln His Met Gln Ala Ala 420 425 430		
	Met Tyr Tyr Ile Leu Thr Arg Gly Trp Ser Asp Asn Gly Asp Pro Met 435 440 445		
20	Phe Asn Gln Thr Trp Ala Thr Phe Ala Met Asn Ile Ala Pro Ala Leu 450 455 460		
	Val Val Asp Ser Ser Cys Leu Ile Met Asn Leu Gln Ile Lys Thr Tyr 465 470 475 480		
25	Gly Gln Gly Ser Gly Asn Ala Ala Thr Phe Ile Asn Asn His Leu Leu 485 490 495		
	Ser Thr Leu Val Leu Asp Gln Trp Asn Leu Met Arg Gln Pro Arg Pro 500 505 510		
30	Asp Ser Glu Glu Phe Lys Ser Ile Glu Asp Lys Leu Gly Ile Asn Phe 515 520 525		
	Lys Ile Glu Arg Ser Ile Asp Asp Ile Arg Gly Lys Leu Arg Gln Leu 530 535 540		
35	Val Leu Leu Ala Gln Pro Gly Tyr Leu Ser Gly Gly Val Glu Pro Glu 545 550 555 560		
	Gln Ser Ser Pro Thr Val Glu Leu Asp Leu Leu Gly Trp Ser Ala Thr 565 570 575		
40	Tyr Ser Lys Asp Leu Gly Ile Tyr Val Pro Val Leu Asp Lys Glu Arg 580 585 590		
	Leu Phe Cys Ser Ala Ala Tyr Pro Lys Gly Val Glu Asn Lys Ser Leu 595 600 605		
	Lys Ser Lys Val Gly Ile Glu Gln Ala Tyr Lys Val Val Arg Tyr Glu 610 615 620		
45	Ala Leu Arg Leu Val Gly Gly Trp Asn Tyr Pro Leu Leu Asn Lys Ala 625 630 635 640		
	Cys Lys Asn Asn Ala Gly Ala Ala Arg Arg His Leu Glu Ala Lys Gly 645 650 655		
50	Phe Pro Leu Asp Glu Phe Leu Ala Glu Trp Ser Glu Leu Ser Glu Phe 660 665 670		
	Gly Glu Ala Phe Glu Gly Phe Asn Ile Lys Leu Thr Val Thr Ser Glu 675 680 685		
55	Ser Leu Ala Glu Leu Asn Lys Pro Val Pro Pro Lys Pro Pro Asn Val		

690

695

700

5 Asn Arg Pro Val Asn Thr Gly Gly Leu Lys Ala Val Ser Asn Ala Leu  
 705 710 715 720

Lys Thr Gly Arg Tyr Arg Asn Glu Ala Gly Leu Ser Gly Leu Val Leu  
 725 730 735

10 Leu Ala Thr Ala Arg Ser Arg Leu Gln Asp Ala Val Lys Ala Lys Ala  
 740 745 750

Glu Ala Glu Lys Leu His Lys Ser Lys Pro Asp Asp Pro Asp Ala Asp  
 755 760 765

15 Trp Phe Glu Arg Ser Glu Thr Leu Ser Asp Leu Leu Glu Lys Ala Asp  
 770 775 780

Ile Ala Ser Lys Val Ala His Ser Ala Leu Val Glu Thr Ser Asp Ala  
 785 790 795 800

20 Leu Glu Ala Val Gln Ser Thr Ser Val Tyr Thr Pro Lys Tyr Pro Glu  
 805 810 815

Val Lys Asn Pro Gln Thr Ala Ser Asn Pro Val Val Gly Leu His Leu  
 820 825 830

25 Pro Ala Lys Arg Ala Thr Gly Val Gln Ala Ala Leu Leu Gly Ala Gly  
 835 840 845

Thr Ser Arg Pro Met Gly Met Glu Ala Pro Thr Arg Ser Lys Asn Ala  
 850 855 860

30 Val Lys Met Ala Lys Arg Arg Gln Arg Gln Lys Glu Ser Arg Gln Gln  
 865 870 875 880

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35 <210> 53  
 <211> 879  
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 <213> Infectious bursal disease virus

40 <220>  
 <221> DOMAIN  
 <222> (1)..(879)  
 <223> /note="Sequence of IBDV D6948-VPl"

45 <400> 53  
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 1 5 10 15

Ala Phe Gly Ile Lys Pro Thr Ala Gly Gln Asp Val Glu Glu Leu Leu  
 20 25 30

50 Ile Pro Lys Val Trp Val Pro Pro Glu Asp Pro Leu Ala Ser Pro Ser  
 35 40 45

Arg Leu Ala Lys Phe Leu Arg Glu Asn Gly Tyr Lys Ile Leu Gln Pro  
 50 55 60

55 Arg Ser Leu Pro Glu Asn Glu Glu Tyr Glu Thr Asp Gln Ile Leu Pro  
 65 70 75 80

5 Asp Leu Ala Trp Met Arg Gln Ile Glu Gly Ala Val Leu Lys Pro Thr  
 85 90 95  
 Leu Ser Leu Pro Ile Gly Asp Gln Glu Tyr Phe Pro Lys Tyr Tyr Pro  
 100 105 110  
 10 Thr His Arg Pro Ser Lys Glu Lys Pro Asn Ala Tyr Pro Pro Asp Ile  
 115 120 125  
 Ala Leu Leu Lys Gln Met Ile Tyr Leu Phe Leu Gln Val Pro Glu Ala  
 130 135 140  
 15 Thr Asp Asn Leu Lys Asp Glu Val Thr Leu Leu Thr Gln Asn Ile Arg  
 145 150 155 160  
 Asp Lys Ala Tyr Gly Ser Gly Thr Tyr Met Gly Gln Ala Thr Arg Leu  
 165 170 175  
 Val Ala Met Lys Glu Val Ala Thr Gly Arg Asn Pro Asn Lys Asp Pro  
 180 185 190  
 20 Leu Lys Leu Gly Tyr Thr Phe Glu Ser Ile Ala Gln Leu Leu Asp Ile  
 195 200 205  
 Thr Leu Pro Val Gly Pro Pro Gly Glu Asp Asp Lys Pro Trp Val Pro  
 210 215 220  
 25 Leu Thr Arg Val Pro Ser Arg Met Leu Val Leu Thr Gly Asp Val Asp  
 225 230 235 240  
 Gly Glu Phe Glu Val Glu Asp Tyr Leu Pro Lys Ile Asn Leu Lys Ser  
 245 250 255  
 30 Ser Ser Gly Leu Pro Tyr Val Gly Arg Thr Lys Gly Glu Thr Ile Gly  
 260 265 270  
 Glu Met Ile Ala Ile Ser Asn Gln Phe Leu Arg Glu Leu Ser Ala Leu  
 275 280 285  
 35 Leu Lys Gln Gly Ala Gly Thr Lys Gly Ser Asn Lys Lys Lys Leu Leu  
 290 295 300  
 Ser Met Leu Ser Asp Tyr Trp Tyr Leu Ser Cys Gly Leu Leu Phe Pro  
 305 310 315 320  
 40 Lys Ala Glu Arg Tyr Asp Lys Ser Thr Trp Leu Thr Lys Thr Arg Asn  
 325 330 335  
 Ile Trp Ser Ala Pro Ser Pro Thr His Leu Met Ile Ser Met Ile Thr  
 340 345 350  
 45 Trp Pro Val Met Ser Asn Ser Pro Asn Asn Val Leu Asn Ile Glu Gly  
 355 360 365  
 Cys Pro Ser Leu Tyr Lys Phe Asn Pro Phe Arg Gly Gly Leu Asn Arg  
 370 375 380  
 50 Ile Val Glu Trp Ile Met Ala Pro Asp Glu Pro Lys Ala Leu Val Tyr  
 385 390 395 400  
 Ala Asp Asn Ile Tyr Ile Val His Ser Asn Thr Trp Tyr Ser Ile Asp  
 405 410 415  
 55 Leu Glu Lys Gly Glu Ala Asn Cys Thr Arg Gln His Met Gln Ala Ala  
 420 425 430

Met Tyr Tyr Ile Leu Thr Arg Gly Trp Ser Asp Asn Gly Asp Pro Met  
 435 440 445  
 5 Phe Asn Gln Thr Trp Ala Thr Phe Ala Met Asn Ile Ala Pro Ala Leu  
 450 455 460  
 Val Val Asp Ser Ser Cys Leu Ile Met Asn Leu Gln Ile Lys Thr Tyr  
 465 470 475 480  
 10 Gly Gln Gly Ser Gly Asn Ala Ala Thr Phe Ile Asn Asn His Leu Leu  
 485 490 495  
 Ser Thr Leu Val Leu Asp Gln Trp Asn Leu Met Lys Gln Pro Ser Pro  
 500 505 510  
 15 Asp Ser Glu Glu Phe Lys Ser Ile Glu Asp Lys Leu Gly Ile Asn Phe  
 515 520 525  
 Lys Ile Glu Arg Ser Ile Asp Asp Ile Arg Gly Lys Leu Arg Gln Leu  
 530 535 540  
 20 Val Pro Leu Ala Gln Pro Gly Tyr Leu Ser Gly Gly Val Glu Pro Glu  
 545 550 555 560  
 Gln Pro Ser Pro Thr Val Glu Leu Asp Leu Leu Gly Trp Ser Ala Thr  
 565 570 575  
 25 Tyr Ser Lys Asp Leu Gly Ile Tyr Val Pro Val Leu Asp Lys Glu Arg  
 580 585 590  
 Leu Phe Cys Ser Ala Ala Tyr Pro Lys Gly Val Glu Asn Lys Ser Leu  
 595 600 605  
 30 Lys Ser Lys Val Gly Ile Glu Gln Ala Tyr Lys Val Val Arg Tyr Glu  
 610 615 620  
 Ala Leu Arg Leu Val Gly Gly Trp Asn Tyr Pro Leu Leu Asn Lys Ala  
 625 630 635 640  
 35 Cys Lys Asn Asn Ala Ser Ala Ala Arg Arg His Leu Glu Ala Lys Gly  
 645 650 655  
 Phe Pro Leu Asp Glu Phe Leu Ala Glu Trp Ser Glu Leu Ser Glu Phe  
 660 665 670  
 40 Gly Glu Ala Phe Glu Gly Phe Asn Ile Lys Leu Thr Val Thr Pro Glu  
 675 680 685  
 Ser Leu Ala Glu Leu Asn Arg Pro Val Pro Pro Lys Pro Pro Asn Val  
 690 695 700  
 45 Asn Arg Pro Val Asn Thr Gly Gly Leu Lys Ala Val Ser Asn Ala Leu  
 705 710 715 720  
 Lys Thr Gly Arg Tyr Arg Asn Glu Ala Gly Leu Ser Gly Leu Val Leu  
 725 730 735  
 50 Leu Ala Thr Ala Arg Ser Arg Leu Gln Asp Ala Val Lys Ala Lys Ala  
 740 745 750  
 Glu Ala Glu Lys Leu His Lys Ser Lys Pro Asp Asp Pro Asp Ala Asp  
 755 760 765  
 55 Trp Phe Glu Arg Ser Glu Thr Leu Ser Asp Leu Leu Glu Lys Ala Asp  
 770 775 780

5 Ile Ala Ser Lys Val Ala His Ser Ala Leu Val Glu Thr Ser Asp Ala  
 785 790 795 800

Leu Glu Ala Val Gln Ser Thr Ser Val Tyr Thr Pro Lys Tyr Pro Glu  
 805 810 815

Val Lys Asn Pro Gln Thr Ala Ser Asn Pro Val Val Gly Leu His Leu  
 820 825 830

10 Pro Ala Lys Arg Ala Thr Gly Val Gln Ala Ala Leu Leu Gly Ala Gly  
 835 840 845

Thr Ser Arg Pro Met Gly Met Glu Ala Pro Thr Arg Ser Lys Asn Ala  
 850 855 860

15 Val Lys Met Ala Lys Arg Arg Gln Arg Gln Lys Glu Ser Arg Gln  
 865 870 875

20 <210> 54

<211> 145

<212> PRT

<213> Infectious bursal disease virus

25 <220>

<221> DOMAIN

<222> (1)..(145)

<223> /note="Consensus sequence of IBDV VP5, whereby X stands for  
 any amino acid"

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Met Val Ser Arg Asp Gln Thr Asn Asp Arg Ser Asp Asp Xaa Pro Ala  
 1 5 10 15

Arg Ser Asn Pro Thr Asp Cys Ser Val His Thr Glu Pro Ser Asp Ala  
 20 25 30

35 Asn Asn Arg Thr Gly Val His Ser Gly Arg His Pro Xaa Glu Ala His  
 35 40 45

Ser Gln Val Arg Asp Leu Asp Leu Gln Phe Asp Cys Gly Gly His Arg  
 50 55 60

40 Val Arg Ala Asn Cys Leu Phe Pro Trp Xaa Pro Trp Leu Asn Cys Gly  
 65 70 75 80

Cys Ser Leu His Thr Ala Glu Gln Trp Glu Leu Gln Val Arg Ser Asp  
 85 90 95

45 Ala Pro Asp Cys Pro Glu Pro Thr Gly Gln Leu Gln Leu Glu Ala  
 100 105 110

Ser Glu Ser Glu Ser His Ser Glu Val Lys His Thr Xaa Trp Trp Arg  
 115 120 125

50 Leu Cys Thr Lys Xaa His His Lys Arg Arg Asp Leu Pro Arg Lys Pro  
 130 135 140

Glu  
 145

55 <210> 55

<211> 145

&lt;212&gt; PRT

&lt;213&gt; Infectious bursal disease virus

&lt;220&gt;

&lt;221&gt; DOMAIN

&lt;222&gt; (1)..(145)

&lt;223&gt; /note="Sequence of IBDV D6948-VP5"

&lt;400&gt; 55

Met Val Ser Arg Asp Gln Thr Asn Asp Arg Ser Asp Asp Glu Pro Ala  
1 5 10 15Arg Ser Asn Pro Thr Asp Cys Ser Val His Thr Glu Pro Ser Asp Ala  
20 25 30Asn Asn Arg Thr Gly Val His Ser Gly Arg His Pro Arg Glu Ala His  
35 40 45Ser Gln Val Arg Asp Leu Asp Leu Gln Phe Asp Cys Gly Gly His Arg  
50 55 60Val Arg Ala Asn Cys Leu Phe Pro Trp Phe Pro Trp Leu Asn Cys Gly  
65 70 75 80Cys Ser Leu His Thr Ala Glu Gln Trp Glu Leu Gln Val Arg Ser Asp  
85 90 95Ala Pro Asp Cys Pro Glu Pro Thr Gly Gln Leu Gln Leu Leu Gln Ala  
100 105 110Ser Glu Ser Glu Ser His Ser Glu Val Lys His Thr Pro Trp Trp Arg  
115 120 125Leu Cys Thr Lys Trp His His Lys Arg Arg Asp Leu Pro Arg Lys Pro  
130 135 140

Glu

145

35

&lt;210&gt; 56

&lt;211&gt; 145

&lt;212&gt; PRT

&lt;213&gt; Infectious bursal disease virus

40

&lt;220&gt;

&lt;221&gt; DOMAIN

&lt;222&gt; (1)..(145)

&lt;223&gt; /note="Sequence of IBDV CEF94-VP5"

&lt;400&gt; 56

Met Val Ser Arg Asp Gln Thr Asn Asp Arg Ser Asp Asp Lys Pro Ala  
1 5 10 15Arg Ser Asn Pro Thr Asp Cys Ser Val His Thr Glu Pro Ser Asp Ala  
20 25 30

50

Asn Asn Arg Thr Gly Val His Ser Gly Arg His Pro Gly Glu Ala His  
35 40 45Ser Gln Val Arg Asp Leu Asp Leu Gln Phe Asp Cys Gly Gly His Arg  
50 55 60

55

Val Arg Ala Asn Cys Leu Phe Pro Trp Ile Pro Trp Leu Asn Cys Gly  
65 70 75 80

Cys Ser Leu His Thr Ala Glu Gln Trp Glu Leu Gln Val Arg Ser Asp  
 85 90 95  
 5 Ala Pro Asp Cys Pro Glu Pro Thr Gly Gln Leu Gln Leu Leu Gln Ala  
 100 105 110  
 Ser Glu Ser Glu Ser His Ser Glu Val Lys His Thr Ser Trp Trp Arg  
 115 120 125  
 10 Leu Cys Thr Lys Arg His His Lys Arg Arg Asp Leu Pro Arg Lys Pro  
 130 135 140  
 15 Glu  
 145

**Claims**

1. An infectious recombinant Infectious Bursal Disease Virus (rIBDV) essentially incapable of growing in a non-bursa-cell or cell derived thereof.
2. An infectious rIBDV having retained at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus (vvIBDV).
3. An rIBDV according to claim 1 having retained at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus (vvIBDV).
4. An rIBDV according to anyone of claims 1 to 3 essentially incapable of growing in a CEF cell, a VERO cell or a QM5 cell.
5. An rIBDV according to anyone of claims 1 to 4 wherein the amino acid sequence of protein VP2 comprises no asparagine at amino acid position 279.
6. An rIBDV according to claim 5 wherein the amino acid sequence of protein VP2 comprises aspartic acid at amino acid position 279
7. An rIBDV according to anyone of claims 1 to 6 wherein the amino acid sequence of protein VP2 comprises no threonine at amino acid position 284.
8. An rIBDV according to claim 7 wherein the amino acid sequence of protein VP2 comprises alanine at amino acid position 284.
9. An rIBDV according to claim 8 wherein the amino acid sequence of protein VP2 at least comprises a stretch of amino acids from about position 279 to 289, preferably from about position 229 to 314, most preferably from about position 214 to 328 as found in a vvIBDV isolate such as shown in Table 8.
10. A method for obtaining an infectious recombinant Infectious Bursal Disease Virus (rIBDV) essentially incapable of growing on a non-bursa-cell derived cell comprising transfecting at least one first cell with a nucleic acid comprising a IBDV genome at least partly derived from IBDV, incubating said first cell in a culture medium, recovering rIBDV from said transfected first cell or said culture medium and propagating said recovered rIBDV in at least one second cell which is permissive for said rIBDV.
11. A method for obtaining an infectious recombinant Infectious Bursal Disease Virus (rIBDV) having retained at least part of the very virulent characteristics of a very virulent Infectious Bursal Disease Virus (vvIBDV) comprising transfecting at least one first cell with a nucleic acid comprising a IBDV genome at least partly derived from a vvIBDV, incubating said first cell in a culture medium, recovering rIBDV from said transfected first cell or said culture medium and propagating said recovered rIBDV in at least one second cell which is permissive for said vvIBDV.

12. A method according to claim 11 or 12 wherein said first cell is a non-bursa-cell derived cell.
13. A method according to anyone of claims 10 to 12 wherein said second cell is a Bursa-cell derived cell.
- 5 14. A method according to anyone of claims 10 to 13 wherein said first cell, such as a CEF cell, a VERO cell or a QM5 cell, is non-permissive for vvIBDV.
- 10 15. A method according to anyone of claims 10 to 14 wherein said first cell has additionally been provided with a helpervirus or a viral protein derived thereof.
16. A method according to claim 15 wherein said viral protein comprises T7-polymerase.
17. A method according to anyone of claims 10 to 16 wherein said rIBDV has at least retained the incapacity to substantially be propagated on a vvIBDV non-permissive cell such as a VERO, a QM5 or CEF cell.
- 15 18. A method according to anyone of claims 10 to 17 wherein said permissive second cell is a primary bursa cell.
19. A method according to anyone of claims 10 to 18 wherein said rIBDV comprises at least a nucleic acid derived from at least a part of genome segment A of vvIBDV.
- 20 20. A method according to claim 19 wherein said nucleic acid encodes at least a functional part of protein VP2.
21. A method according to anyone of claims 10 to 20 wherein said rIBDV comprises at least a nucleic acid derived from a serotype II IBDV.
- 25 22. A method according to anyone of claims 10 to 21 wherein said rIBDV is lacking at least one immunodominant epitope specific for a serotype I IBDV.
23. An infectious mosaic IBDV (mIBDV) comprising a rIBDV wherein at least one genome segment comprises nucleic acid derived from at least two different Birna virus isolates.
- 30 24. A mIBDV according to claim 23 wherein at least one of said isolates is a wIBDV.
- 25 25. A mIBDV according to claim 23 or 24 characterised by its incapacity to substantially be propagated on a wIBDV non-permissive cell such as a VERO, a QM5 or CEF cell.
26. A mIBDV according to anyone of claims 23 to 25 characterised by its capacity to substantially be propagated on a vvIBDV permissive cell such as a primary bursa cell.
- 40 27. A mIBDV according to anyone of claims 23 to 26 wherein at least one of said isolates is a serotype II IBDV.
28. A mIBDV according to anyone of claims 23 to 27 lacking at least one immunodominant epitope specific for a serotype I IBDV.
- 45 29. A vaccine comprising a rIBDV according to anyone of claims 1 to 9 or 23 to 28.

50

55

Figure 1: antibody titer in broilers (actively vaccinated with intermediate vaccine at 3 weeks)

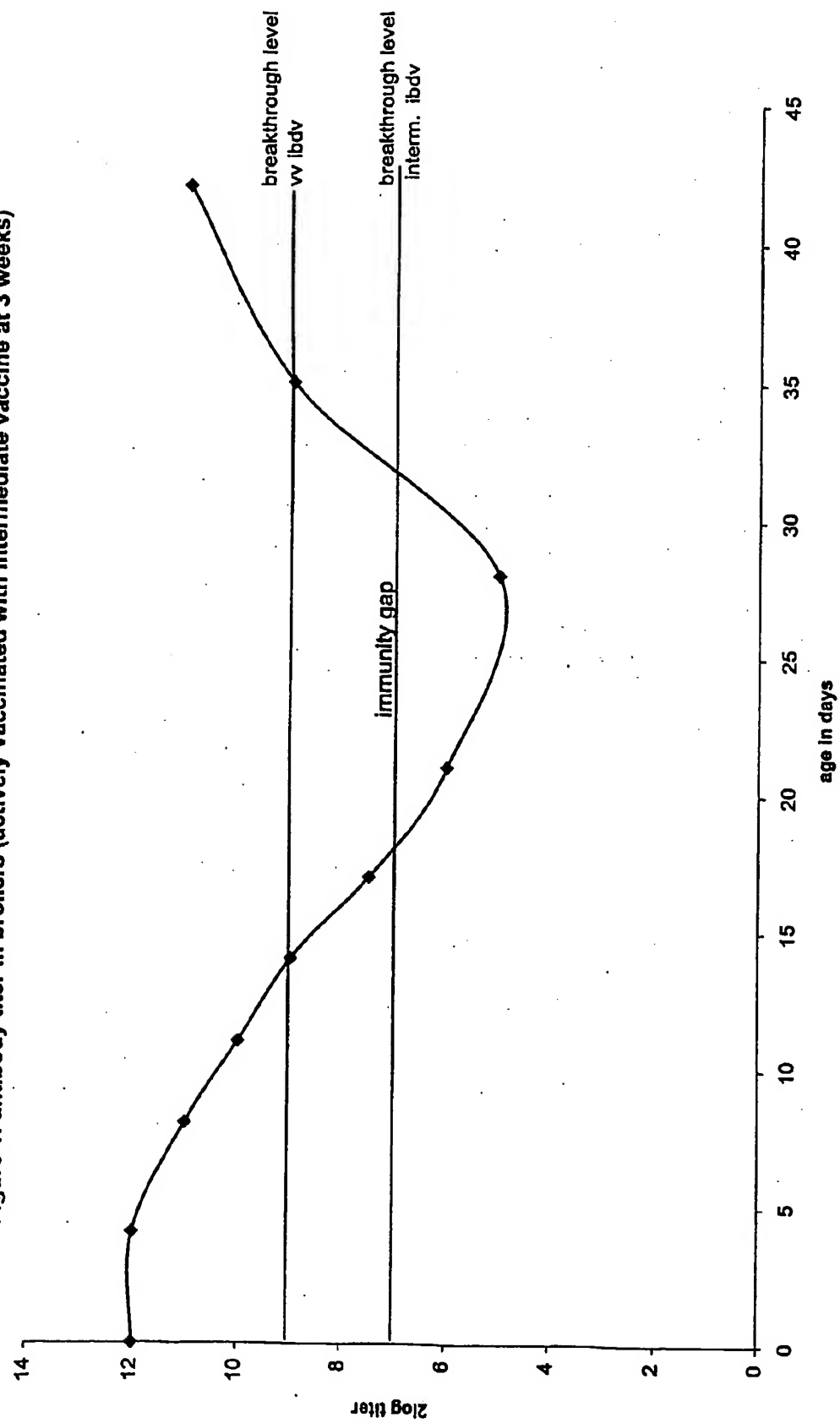


Fig. 2a Alignment of IBDV A-segment cDNA sequences

Consensus	GGATACGATC GGCTCTGACCC CGGGGGAGTC ACCCGGGGAC AGGCYGMCAA GGYCTTGTC CAGGATGGAA CTCCCT	75
CEP94-A	.....	.....
D6948-A	.....	.....
TY89-A	.....	.....
Consensus	CCTTCTACAA YGCTATCATT GATGGTYAGT AGAGATCAGA CAAACGATCG CAGCGATGAC AACCTGCAA GATCA	150
CEP94-A	..... C..... C.....	A.....
D6948-A	..... T..... T.....	G.....
TY89-A	.....	.....
Consensus	AACCCAAACAG ATTGTTCCGT TCATACGGAG CCTTCTGATG CCACCAACCG GACCGGGCTC CATTCCGGAC GACAC	225
CEP94-A	.....	.....
D6948-A	.....	.....
TY89-A	.....	.....
Consensus	CCTRGAGAAG CACACTCTCA GGTCAGAGAC CTCGACCTAC AATTTGACTG TGGGGGACAC AGGOTCAGGG CTAAT	300
CEP94-A	..... G.....	.....
D6948-A	..... A.....	.....
TY89-A	.....	.....
Consensus	TGTCTTTTC CCTGGWTTCC CTGGCTCAAT TGTGGGTGCT CACTACACAC TGCAGAGCAA TGGGAACATC AAGTT	375
CEP94-A	.....	.....
D6948-A	..... A.....	.....
TY89-A	..... T.....	.....
Consensus	CGATCAGATG CTCCCTGACTG CCCAGAACCT ACCGGCCAGY TACAACACTA GCAGGCTAGT GAGTCGGAGT CTCAC	450
CEP94-A	.....	.....
D6948-A	.....	.....
TY89-A	.....	.....
Consensus	AGTGGAGGTCA AGCACACTYC CTGGTGGCGT TTATGCACTA AAYGGCACCA TAAACGCCGT GACCTTCCAA GGAAG	525
CEP94-A	.....	.....
D6948-A	..... T.....	.....
TY89-A	..... C.....	.....
Consensus	CCTGAGTGAA CTGACAGATG TEAGCTACAA TGGGTTGATG TCTGCAACAG CCAACRTCAA CGACAAAATY GGGAA	600
CEP94-A	.....	.....
D6948-A	.....	.....
TY89-A	.....	.....
Consensus	CGTCCTAAGTA GGGGAAGGGG TAAACCTCTT CAGCTTACCC ACATCATATG ATCTTGGGT AATGAGRCYT GGTGA	675
CEP94-A	.....	.....
D6948-A	..... C.....	..... G..... T.....
TY89-A	..... A.....	..... A..... C.....
Consensus	CCCCATTCCC GCWATAGGGC TYGACCCCCA AATGGTACCH ACATGAGACA GCACTGACAG GCCAGAGTC TACAC	750
CEP94-A	..... A..... T.....	..... C.....
D6948-A	..... T..... C.....	..... A.....
TY89-A	.....	.....
Consensus	CATAACTGCA GCGGATGATT ACCAATTCTC ATCACAGTAC CAASCAAGTG GGGTAACAAAT CACACTGTT TCAGC	825
CEP94-A	.....	.....
D6948-A	.....	.....
TY89-A	.....	.....
Consensus	YAAATTTGAT CCCTTCACAA CCCTCAAGT TGGGGGAGAR CTCUTGTTTC AACACAGCGT CCAGGGCTT RTACT	900
CEP94-A	..... C..... T.....	..... G..... T.....
D6948-A	..... T..... C.....	..... A..... C.....
TY89-A	.....	.....
Consensus	GGGGYGYACC ATCTTACCTYA TAGGCTTTGA TGGGACWGCG GTAATCACCA GRCGCTGTGGC CGCARACAAAT GGGCT	975
CEP94-A	..... C..... C.....	..... A..... G.....
D6948-A	..... T..... T.....	..... T..... A.....
TY89-A	.....	.....

Fig. 2a Alignment of IBDV A-segment cDNA sequences

Consensus	RACGRCCGGC ACYGACAAAC C TTWIGCCATT CAATTTGTG ATTCCAACMA RCGAGATAAC CCAGCCANTC ACATC	1050
CEP94-A	G...A.....C.....T.....C.....A.....A.....	1050
D6948-A	A...G.....T.....A.....A.....C.....G.....	1050
TY89-A	-----	-----
Consensus	CATCAAATG GAGATAGTGA CCTCCAAAAG TGGTGGTCAG GCRGGGGATC AGAATGCTG GTCRGCAGW GGGAG	1125
CEP94-A	.....A.....G.....G.....A.....	1125
D6948-A	.....G.....A.....A.....T.....	1125
TY89-A	-----	-----
Consensus	CCTAGCAGTG ACCATCCAYG GTGGCAACTA TCCAGGGGCC CTCCGTCCCG TCACRCTAGT RGCCCTACGAA AGAGT	1200
CEP94-A	.....T.....G.....G.....A.....	1200
D6948-A	.....C.....A.....A.....	1200
TY89-A	-----	-----
Consensus	GCCAAACAGGA TCGTGTGTTA CGGTGCGYGG GGTGAGCAAC TTGAGCTGA TCCCRAATCC TGAACTAGCA AAGAA	1275
CEP94-A	.....C.....T.....	1275
D6948-A	.....T.....C.....	1275
TY89-A	-----	-----
Consensus	CCTGGTYACA GAATAACGGCC GATTGACCC AGGAGCCATG AACTACACAA ATTGATACT GAGTGAGAGG GACCG	1350
CEP94-A	.....T.....	1350
D6948-A	.....C.....	1350
TY89-A	-----	-----
Consensus	TCTTGGCATE AAQACCGTNT GGCCAAACAAAG GGAGTACACT GACTTTGCGY ARTACTTCAT GGAGGTGGCC GACCT	1425
CEP94-A	.....C.....T.....A.....	1425
D6948-A	.....A.....C.....G.....	1425
TY89-A	-----	-----
Consensus	CAACTCTCCC CTGAAGATTG CAGGAGCATT YGGCTTCAAA GACATAATCC GGGCCTTAAG GAGGATAGCT GTGCC	1500
CEP94-A	.....C.....A.....	1500
D6948-A	.....T.....C.....	1500
TY89-A	-----	-----
Consensus	GGTGGTCTCY ACAYTGTTC CACCGGCCGTC TCCCTAGCC CATCCAATTG GGGAGGGT AGACTACCTG CTGGG	1575
CEP94-A	.....C...T.....T.....	1575
D6948-A	.....T...C.....C.....	1575
TY89-A	-----	-----
Consensus	CGATGAGGCA CAGGCTGCTT CAGGAACATGC TCGAGCCCG TCAGGAAAG CAAGAGCTGC CTCAGGCCGC ATARG	1650
CEP94-A	.....	1650
D6948-A	.....	1650
TY89-A	-----	-----
Consensus	GCAGCTRACT CTCGCCCCCG ACAAGGGTA CGAGGTACTC GCGAAATCTT TYCAGGTGCC CCAGAAATCCY GTAGT	1725
CEP94-A	.....G.....A.....C.....C.....	1725
D6948-A	.....A.....G.....T.....T.....	1725
TY89-A	-----	-----
Consensus	CGACGGGATT CTYGGTTCAC CTGGGTTACT CGCGCGYGA CACAACCTCG ACTGGTGTGTT RAGAGAGGGT GCCAC	1800
CEP94-A	.....T.....G.....T.....A.....	1800
D6948-A	.....C.....A.....C.....G.....	1800
TY89-A	-----	-----
Consensus	GCTATGCCCT GTGGTYATYA CGACAGTGGG AGAYGCCATG ACACCCAAAG CAYTGAACAG CAAATGTTT GCTGT	1875
CEP94-A	.....T...T.....C.....T.....	1875
D6948-A	.....C...C.....T.....C.....	1875
TY89-A	-----	-----
Consensus	CATTGAAGGC GTGCGAGAAG AYCTCCAACC TCCWCTCAAAAGGAGATCT TCATACGAAAC TCTCTCYGGA CAYAG	1950
CEP94-A	.....C.....T.....T.....T.....C.....C.....	1950
D6948-A	.....T.....A.....C.....C.....T.....	1950
TY89-A	-----	-----

Fig. 2a Alignment of IBDV A-segment cDNA sequences

Consensus	AGTCTATGGA TATGCTCCAG ATGGGTTACT TCCACTGGAG ACTGGGAGAG AYTACACCGT KGTOCCAATA GATGA	2025
CEF94-A	.....	.....C.....T.....
D6948-A	.....	.....T.....G.....
TY89-A	.....	.....
Consensus	TGTCCTGGAC GACAGCATTG TGCTGTCCAA AGAYCCCATG CCTCCCTATG TGGGAAACAG YGGAAAYCTA GCCAT	2100
CEF94-A	.....	.....T.....T.....
D6948-A	.....	.....C.....C.....
TY89-A	.....	.....
Consensus	AGCTTACATG GATGTGTTTC GACCCAAAGT CCCMATCCAT GTGGCYATGA CGGGAGCCCT CAAYGCYTRT GGCGA	2175
CEF94-A	.....	.....A.....T.....
D6948-A	.....	.....C.....C.....
TY89-A	.....	.....
Consensus	GATTGAGAAAGT GTRAGCTTTA GAAGCACCAA GCTGCCACT GCACACCGAC TTGGCCTYAA GTTGGCTGGT CCCGG	2250
CEF94-A	.....A.....A.....	.....T.....
D6948-A	.....C.....G.....	.....C.....
TY89-A	.....	.....
Consensus	WGCATTGAY GTRAAACACCG GGYCCAACGT GGCRACGTTT ATCAAAACGTT TYCCTCACAA TCCMCGMGAC TGGGA	2325
CEF94-A	.....A.....C.....T.....A.....	.....C.....A.....C.....
D6948-A	.....T.....T.....C.....G.....	.....T.....A.....C.....
TY89-A	.....	.....C.....C.....A.....
Consensus	CAGGYTCCTY TACCTCAACC TWCCTTAYCT YCCACCHAAW GCGNGGACCGY AGTWCMSAYCT KGCGCTGGCH GCHTC	2400
CEF94-A	.....C.....C.....	.....A.....A.....C.....
D6948-A	.....C.....T.....	.....T.....A.....C.....
TY89-A	.....T.....A.....C.....	.....A.....G.....A.....C.....T.....
Consensus	NGAGTTCAA GAGACCCCG AACTCGARRR YGCGTSHGW GCMATGGAMG CGCGCGHAA CGTSGACCCA YTTRT	2475
CEF94-A	.....A.....	.....G.....A.....
D6948-A	.....A.....C.....	.....A.....C.....
TY89-A	.....C.....A.....	.....G.....C.....
Consensus	CCRMTCWGCD CTCMRBGST TCTATGTGGYT GGAAGARAAY GGGATTGTRA CYGAYATGCC YAACTTCGCM CTCAG	2550
CEF94-A	.....AA.....T.....A.....	.....AGT.....G.....
D6948-A	.....AA.....T.....G.....	.....AGC.....G.....
TY89-A	.....GC.....A.....T.....	.....CAG.....C.....
Consensus	CGACCCGAAC GCHCAYHGGG TGKRMIAATT YCTHGCCAAAY GCNCMCMARG CHGGHAGCAA GTGCGCARAGR GCCAA	2625
CEF94-A	.....C.....TC.....	.....CGA.....T.....T.....C.....
D6948-A	.....C.....TC.....	.....CGC.....T.....C.....
TY89-A	.....A.....CA.....	.....AAA.....C.....A.....T.....T.....C.....G.....C.....A.....
Consensus	GTTAYGGGACR GCWGCGTACG GAOTGGAGGC YMGRGGGCCCT ACDCGAGARG ARGCACAGAG GGARAAAGAC ACACG	2700
CEF94-A	.....C.....G.....A.....	.....A.....TC.....
D6948-A	.....C.....G.....A.....	.....CCG.....T.....G.....A.....
TY89-A	.....T.....C.....G.....T.....	.....TA.....A.....G.....A.....C.....
Consensus	GATCTCMAAG AAGATGGARA CBATGGGCAT CTACTTGYCA ACACCRGAAT GGCTTGGCACT CAAYGGGCAC CGRGG	2775
CEF94-A	.....A.....	.....G.....C.....
D6948-A	.....A.....	.....G.....T.....
TY89-A	.....C.....	.....A.....G.....
Consensus	SCCAAGCCCC GGCCAGCTVA AGTACTGGCA RAACACAMGA GAAATACCDG AHCCMAACGA GGACTTAYCYA GACTA	2850
CEF94-A	.....G.....	.....A.....G.....C.....
D6948-A	.....G.....	.....G.....C.....
TY89-A	.....C.....	.....A.....A.....
Consensus	YGTGCGAYGCR GAGAAGAGCC GGTTGGCRTC AGAAGAACAR RTCYTAAGGG CAGCYACGTC GATCTACGGG GCTCC	2925
CEF94-A	.....C.....T.....A.....	.....A.....A.....C.....
D6948-A	.....C.....T.....A.....	.....A.....A.....C.....
TY89-A	.....T.....C.....G.....	.....G.....G.....T.....

Fig. 2a Alignment of IBDV A-segment cDNA sequences

Consensus	AGGACAGGCW GARCCACCCC AAGCTTTCAT AGACGAAGTY GCCARRGTCT ATGAAATCAA CCATGGRCGT GGYCC	3000
CEP94-A	.....A ..G..... .T..... .T.....A..... .A..... .C..	3000
D6948-A	.....A ..G..... .C..... .C.....A..... .G..... .C..	3000
TY89-A	.....T ..A..... .C.....C.....G.....G.....T..	704
Consensus	MAACCARGAR CAGATGAARG AYCTGCTCYT GACTGGATG GAGATGAAGC ATCGCAATCC CAGGGGGCT CYACC	3075
CEP94-A	A.....A ..A ..A ..T.....T..... .T..... .T..	3075
D6948-A	C.....A ..A ..A ..T.....C..... .C..... .C..	3075
TY89-A	A.....G ..G ..G ..C.....C..... .C..... .C..	779
Consensus	AAAGCCHAAG CCAAAACCCA ATGCTCCANC ACAGAGACCC CCTGGWCGGC TGGGCCGCTG GATCAGGRGB GTCTC	3150
CEP94-A	.....C..... .A..... .A..... .T..... .A.C..... .A.C.....	3150
D6948-A	.....C..... .A..... .A..... .T..... .G.T..... .G.T.....	3150
TY89-A	.....A..... .T..... .T..... .A..... .A.G..... .A.G.....	854
Consensus	TGAYGAGGAC YTKGAGTGAG GYNCCCTGGGA GTCTCCCGAC ACCACCCGCG CAGGYGTGGA CACCAATTMR KMMHT	3225
CEP94-A	...T..... C.T..... TA..... .T..... .T..... CG GACT.	3225
D6948-A	...T..... C.T..... CT..... .C..... .C..... CG GCCA.	3225
TY89-A	...C..... T.G..... CT..... .T..... .AA TCAC.	929
Consensus	ASWRMAYCS AATTTGGATC CGTTCGGGG TCCCC	3260
CEP94-A	.CAAC..C.C .....	3260
D6948-A	.CAAC..C.C .....	3260
TY89-A	.GTGA..T.G .....	964

Fig. 2b Alignment of IBDV B-segment cDNA sequences

Consensus	GGATACGATG GGTCTGACCC TCTGGGAGTC AGCAATTAC GTCGGCTACTA GGGGYGATRM CCRCCGCTRG CTGCC	75
CEF94-B	.....	.....C..AA..G..G..
D6948-B	.....	.....T..GC..A..A..
Consensus	ACGTTAGTGG CTCCCTCTCT TGATGATTCT RCCACCATGA GTGACRTTTT CAAYAGTCCA CAGGGCGGAA GCANG	150
CEF94-B	.....	.....G.....A.....C.....
D6948-B	.....	.....A.....G.....T.....
Consensus	ATATTCAGCAG CGTTGGCAT AAAGCCTACW GCTGGACARG AYGTGGAGA ACTCTTGATC CCTAARGTTT CGCTG	225
CEF94-B	.....C.....	.....T.....A..C.....T.....A..T.....
D6948-B	.....A.....	.....A.....G.....T.....C.....G..C.....
Consensus	CCACCTGAGG ATCCSYTKGC CAGCCCTAGT CGWCTGGCHW AGTTCCTCAQ RGARAACCGC TACAARRTTY TGCAG	300
CEF94-B	.....GC..T.....	.....A..A.....A..G.....AG..T.....
D6948-B	.....CT..G.....	.....T..C.....G..A.....GA..C.....
Consensus	CCACGGTCTC TCCCCYAGAA TGAGGGAGTAT GAGACCGAYC AAATACTCCC WGACYTTAGCW TGGATGMGRG AGATA	375
CEF94-B	.....G..C.....	.....C.....A..T..A.....C..A.....
D6948-B	.....A..T.....	.....T.....T..C..T.....A..G.....
Consensus	GARGGRGCTG TTTTAAACCC MACYCTATCT CTCCCCYATTG GAGAYCAGGA GTACTTCCW AARTACTACC CAACA	450
CEF94-B	.....A..G.....	.....C..T.....T..T.....A..G.....
D6948-B	.....G..A.....	.....A..C.....C.....T..A.....
Consensus	CAYCGCCCKA GCAAGGARAA GCCCAATGGG TACCCGGCGM AYATCGCAYT ACTCAAGCAG ATGATYTAZY TGTPT	525
CEF94-B	.....T.....T.....G.....	.....A..C.....C.....T..C.....
D6948-B	.....C.....G.....	.....A..C.....C..T.....T.....C..T.....
Consensus	CTCCAGGTTC CNGAGGCCAM NGAKRRCCTW AARGATGARG TAAACCTTWT RACCCAAAAC ATWAGRGAAY ARGCC	600
CEF94-B	.....A.....A..C..GGG..A..G..A..A..A..CT..G.....	.....A..G..C..G..A..T..A..T..A..
D6948-B	.....C.....C..A..TAA..T..A..G..C..A..AC..A.....	.....T..A..T..A..T..A..
Consensus	TAYGGRAGTG GGACCTACAT GGGACARGCM ACYNGACTTG TKGCCYATGAA RGAGGTGTYCC ACTGGRAGAA ACCCA	675
CEF94-B	.....T..A.....A..A..TC.....G..C.....G..C.....A.....	.....A..G..C..G..A..T..A..T..A..
D6948-B	.....C..G.....G..C..CA.....T..T..A.....T..A.....G..	.....A..C.....A..T..A..G..
Consensus	AACAARGATC CTCTAAAGCT TGGGTACACY TTTGAGAGCA TNGCSCAGCT ACTTGACATC ACWYTACCGG TAGGC	750
CEF94-B	.....G.....	.....T.....C..G.....A..C.....
D6948-B	.....A.....	.....C.....A..C.....TT.....
Consensus	CCACCCGGTG AGGATGACAA GCCCTGGGTR CCACTCACAA GRTGCGGTC AMGGATGTTG GTWCTGACGG GMGAC	825
CEF94-B	.....G.....	.....A.....C.....A.....A.....A..A..A..
D6948-B	.....A.....	.....G.....A.....A.....T..C..C..
Consensus	GTAGATGGSG ANTTTGAGGT TGARGAYTAC CTTCCCCAAA TCAACCTCAA GTCATCAAGT GGACTRCMT ATGTM	900
CEF94-B	.....C..C.....A..T..	.....A..A..A..A..A..
D6948-B	.....G..A.....G..C..	.....G..C..G..C..T..
Consensus	GGTCGACCCA AAGGAGARAC WATGGGGAG ATGATAGCYA TNCRAACCA GTTCTCTYGA GAGCTATCAR CRCTG	975
CEF94-B	.....G..A..C..T..G..	.....T..C..A..A..CA..A..A..A..
D6948-B	.....A..T..G..C..	.....C..A..G..C..TC..G..G..
Consensus	YTGAAGCARG GTGCGAGGAC AAARGGCTTAC AACAGAAGA AGCTTCTTCAG CATGTYAAGT GACTAYTGGT ACTTA	1050
CEF94-B	.....T..A.....G..A.....A..A.....T..A.....T..T..	.....T..A..T..A..T..T..
D6948-B	.....C..G.....A..G.....G..G.....C..C..C..	.....G..C..G..C..T..C..C..
Consensus	TCATGYYGGC TTTTGTTTCC MAAGGCTGAR AGGTACCGACA AAAGYACATG GCTCACCAAG ACCCGKAACA TATGG	1125
CEF94-B	.....C.....A.....A.....T..	.....G..G..G..T..
D6948-B	.....T.....C.....G.....C..C..	.....T..T..T..T..
Consensus	TCAGCTCCAT CNGAACACAA CCTCATGATC TCAWTGATHA CCTGGCCCGT GATGTCCAAY AGCCCAAAYA ACCTG	1200
CEF94-B	.....C.....	.....T..C.....C..T..
D6948-B	.....A.....A..A..A..A..A..	.....T..C..C..T..C..

Fig. 2b Alignment of IBDV B-segment cDNA sequences

Consensus	TTGAAACATTG ARGGGTGTCC RTCACTCTAC AATTCACCC CTTTACAGGG WGGGUTRAAC AGGATCCTSG ACTGG	1275
CEP94-B	.....A.....A.....A.....C.....A..T.G.....C.....	1275
D6948-B	.....G.....G.....G.....T.....T..C.A.....G.....	1275
Consensus	ATATGGCYC COGANGAACCC CAAGGCTTWW GTATATGCCG ACRACATATA CATTOTYAC CTCACACAGT GGTAC	1350
CEP94-B	...T...C...A.....TC.T.....G.....C...A.....	1350
D6948-B	...A...T...T.....CT.A.....T.....T...C.....	1350
Consensus	TCAATTGACC TAGAGAAAGG TGAGGCAAC TGCACTCGYC AARACATCA CCGCCGATG TACTACATTC TYACC	1425
CEP94-B	.....T.....T.C.....A.....A.....A.....A.....	1425
D6948-B	.....G.....G.T.....G.....C.....C.....T.....	1425
Consensus	AGAGGTTGCT CGAGAACCGG YGACCCCHATG TTCATTCARA CATGGGCCAC CTTTGCATG AACATTGCCG CGCT	1500
CEP94-B	....G....A..C....C....A....A....C....T....	1500
D6948-B	....A....C..T....T....C....G....G....A....	1500
Consensus	CTAGTKGTGG ACTCATCRTG YCTGATWAG AACCTKCARA TYAAGACHTA TGGTCAAGGC AGYGGGAATG CAGCC	1575
CEP94-B	....G....G..C....A....G..A..T....C....C....	1575
D6948-B	....T....A..T....T....T..G..C....A....T....	1575
Consensus	ACSTTCATCA ACAACCAYCT YYTRAGCAAC CTWGTGCTWG ACCAGTGGAA CYTGATGARR CARCCYAGWC CAGAC	1650
CEP94-B	...G.....C..CT.G....G..A..T.....C.....GA..G..C..A.....	1650
D6948-B	...C.....T..TC.T....C..T..A.....T.....AG..A..T..T.....	1650
Consensus	AGCGGARGAGT TCAARTCAAT TGARGACAACT CTGGYATCA ACTTYAAGAT TGAGAGGTCC ATTGTGAYA TYAGG	1725
CEP94-B	....G....A.....G.....A..T.....T.....T.....C..T....	1725
D6948-B	....A....G.....A.....G..C.....C.....C.....C..T....	1725
Consensus	GGCAAGCTSA GACAGCTTGT CCYCCCTGCA CAACCAAGGT ACCTGAGTGG RGGGGTTGAA CCAGARCAAY CCAGC	1800
CEP94-B	....G.....T.....G.....T.....A..T.....A.....T.....	1800
D6948-B	....C.....C.....A.....C.....A.....G..C.....	1800
Consensus	CCAACCTGNG AGCTKGACCT ACTHGGRTGG TCGGCAACWT ACAGCAAAGA TCTYGGATC TATGTGCCGG TGCTT	1875
CEP94-B	....T....T....A..G....A..T..A.....C.....	1875
D6948-B	....A....G....C..A....T..A..T.....T.....	1875
Consensus	GACAAAGAAC GCTTATTTG YTCGTGCG TATCCAARG GRGTAGAGAA YAARAGYCTC AARTCCAARG TYGGG	1950
CEP94-B	.....C.....T.....G..A.....C..G..T....G.....A..T....	1950
D6948-B	.....T.....C.....A..G.....T..A..C....A....G..T....	1950
Consensus	ATCGGACG CATAACAAGT WGTCAGGTAY GAGGGCTGTA GGTGGTAGG TGGTGGAAAC TACCCACTCC TGAC	2025
CEP94-B	.....G.....G..A.....T.....T.....T.....	2025
D6948-B	.....A.....A..T.....C.....T.....	2025
Consensus	ANACCTYCA AGAAAYAYGC ARGYGCGGT CGGGGGCATC TGGAGGCCAA GGGGTCCCR CTGGAYGAAT TCTH	2100
CEP94-B	....C.....T..C..G..C.....A.....A.....A.....A.....	2100
D6948-B	....T.....C..T..A..T..A.....G.....T.....C.....	2100
Consensus	GCCGAGTGCTT CGAGAGYTCG MGAGTTGGW GARGCYTTG AAGGCTTCAA YATCAAGCTG AGCTAACAY CEGAG	2175
CEP94-B	.....T..C....A.....T..G..C.....T.....C.....T..T....	2175
D6948-B	.....A..T....C.....A..A..T.....C.....A....C..G....	2175
Consensus	AGCCCTGGCG AACTKAACAR RCCAGTACCC CCCAACCCYC CAAATGCAA CAGACCAAGTC AACACYGGKG GRCTH	2250
CEP94-B	....A....G....A..G.....G..C.....T..G..A..C.....	2250
D6948-B	....C.....T..G..A.....A..T.....C..T..G..A.....C..T..G..A.....	2250
Consensus	AAGGCAGTC ACAAAGCCCT CAAGACCGG CGGTAYAGRA AYGAAGCCGG ACTRAGTGGY CTCGTCTTC TAGCC	2325
CEP94-B	.....C.....T.....C..G..C.....G.....T.....T.....	2325
D6948-B	.....T.....C.....C..T..A..T.....A..C.....C.....C..T.....	2325
Consensus	ACGCGMNGMA GCGGWCTRCA RGAYGCAGTY AAGGCCAAGG CAGARGCCGA GAAACTCCAC AAGTCYAAGC CGHAY	2400
CEP94-B	...A..AA.A....T..G..A..T..T.....A.....C.....A..C.....	2400
D6948-B	...C..CC.C....A..A..G..C..C.....G.....T....C..T.....	2400

Fig. 2b Alignment of IBDV B-segment cDNA sequences

Consensus	GACCCCGATG CAGACTGGTT YGAAGRTCA GAAACYCTGT CAGACCTKCT GGAGAAAGCC GACATYGCCA GCAAG	2475
CEF94-B	..... C...A.A... ....T.....T.....C.....	2475
D6948-B	..... T...C.G... ....C.....G.....T.....	2475
Consensus	GTCGGCYCACT CAGCAGCTGGT GGAACAAAGC GACGGCYCTTG AAGCRGTYCA GTCRACYTCN GTGTAACACYC CAAAG	2550
CEF94-B	.....C..... ....C.....A..T.. ....G..T..C.....T..C..	2550
D6948-B	.....T..... ....T.....G..C.. ....A..C..A.....C..A..	2550
Consensus	TACCCAGARG TYAAGAAACCC ACAGACCCGCC TCCAACCCCG TTGTTGGGCT CCACCTGCC GCGAAGAGRG CCACC	2625
CEF94-B	.....A..C..... ....A.....	2625
D6948-B	.....G..T..... ....G.....	2625
Consensus	GGTGTCCAGG CGGCTCTTCT CGGAGCAGGR ACGAGCAGAC CAATGGGGAT GGAGGGYCCA ACACGGTCCA AGAAC	2700
CEF94-B	.....C..... ....A.....C.....	2700
D6948-B	.....A.....G.....T.....	2700
Consensus	GCCGTGAAA TGGCCAAAMG GCGGCCAACGC CAAARGAGA GCGGCCAYA GCCATGATGG GAACCACTCA AGAAG	2775
CEF94-B	.....C..... ....G.....C.....	2775
D6948-B	.....A.....A.....T.....	2775
Consensus	AGGACACTAA YCCCAGACCC CGTATCCCCG GCCTTCGCGCT GCGGGGGCCC CC	2827
CEF94-B	.....T..... ....	2827
D6948-B	.....C..... ....	2827

Fig. 3a IBDV polyprotein alignment

Consensus	MTNLQDQTOQ IVPPFIRSLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIVPF PGPPGSIVGA HYTLQ	75
CEF94-PP	.....	75
D6948-PP	.....	75
TY89-PP	.....	
Consensus	SNGNYKFDQN LLTAGNLPAS YNYCRLVRSR LTVRSSTLPG GYVALNGTIN AVTPQGSSLB LTDWSYNGLM SATAN	150
CEF94-PP	.....	150
D6948-PP	.....	150
TY89-PP	.....	
Consensus	INDXIGNVLV GCGUTVILSLP TSYDLGTVRL GDPPIAIGLD PKMVATCDSR DRPKVYTITA ADDYQFSSQY Q.GGV	225
CEF94-PP	.....	225
D6948-PP	.....	225
TY89-PP	.....	
Consensus	TITLPSANID AITSLS.GGE LVFQTSV.GL .LGATIYLIG FDGTAVITRA VAA.NGLT.G TDNL.PFN.V IPT.E	300
CEF94-PP	.....V.....H.....V.....N.....T.....L.....N.	300
D6948-PP	.....I.....Q.....I.....D.....A.....M.....I.....S.	300
TY89-PP	.....	
Consensus	ITQPITSIKL EIVTSKSGGQ AGDQMSWSA. GSLAVTIHGG NYPGALRPVT LVAYERVATG SVVTVAGVSN FELIP	375
CEF94-PP	.....R.....	375
D6948-PP	.....S.....	375
TY89-PP	.....	
Consensus	NPELAKNLVT EYGRFDPGAM NYTKLILSER DRIGIKTVWP TREYTDPREY FMEVADLNSP LKIAAGAPGPK DIIRA	450
CEF94-PP	.....	450
D6948-PP	.....	450
TY89-PP	.....	
Consensus	.RRIAVPVVS TLPPPAAPLA HAIGEGVDYL LGDEAQAAASG TARAASGKAR AASGRIRQLT LAADKGYEVV ANLFQ	525
CEF94-PP	I.....	525
D6948-PP	L.....	525
TY89-PP	.....	
Consensus	VPQNPVVVDGI LASPG.LRGA HNLDCLVREG ATLFFFVITT VEDAMTPKAL NSKMPAVIEG VREDLQPPSQ RGSPF	600
CEF94-PP	.....V.....	600
D6948-PP	.....I.....	600
TY89-PP	.....	
Consensus	RTLSGHRVYG YAPDGVLPLE TGRDUTTVVPI DDVMDDSIML SKDPIPPIVG NSGNLAIAYM DVPRPKVPIH VAMTG	675
CEF94-PP	.....	675
D6948-PP	.....	675
TY89-PP	.....	
Consensus	AIINA.GEIE. VSPRSTKLAT AHRIGLKLKG PGAFDVNTG. NWATFIKRPP HNPRDWDRLP YLNLPYLPPN AGRQY	750
CEF94-PP	....C....K.....P.....	750
D6948-PP	....Y....N.....S.....	750
TY89-PP	.....T.....F.....	28
Consensus	HLAMAASEPK ETPELESAVR AMEAAANVDP LFQALSALSPM WLEENGIVTD MANFALSDPN AHRMRNPLAN APQAG	825
CEF94-PP	.....	825
D6948-PP	D.....	825
TY89-PP	....L.....D....R...Q....K.....	103
Consensus	SKSQRAXYGT AGYGVEARGP TPSEAAQREKD TRISKRMETH GIYPATPEWV ALNGHRGPSP GQLKYWQNTRE EIPDP	900
CEF94-PP	.....	900
D6948-PP	.....	900
TY89-PP	.....E.....	178
Consensus	NEDYLDYVHA EKSRLASEEQ ILRAATSIYG APGQARPPQA FIDEVAKVYE INHGRGPQG QMKDLLLTA MNGCHR	975
CEF94-PP	.....	975
D6948-PP	.....	975
TY89-PP	....P.....V.....R.....	253
Consensus	NPRRAPPKPK PKPNAPTORP PGRLGRWIRT VSDEDLE	1012
CEF94-PP	....L.....	1012
D6948-PP	.....A.....	1012
TY89-PP	....S.....	290

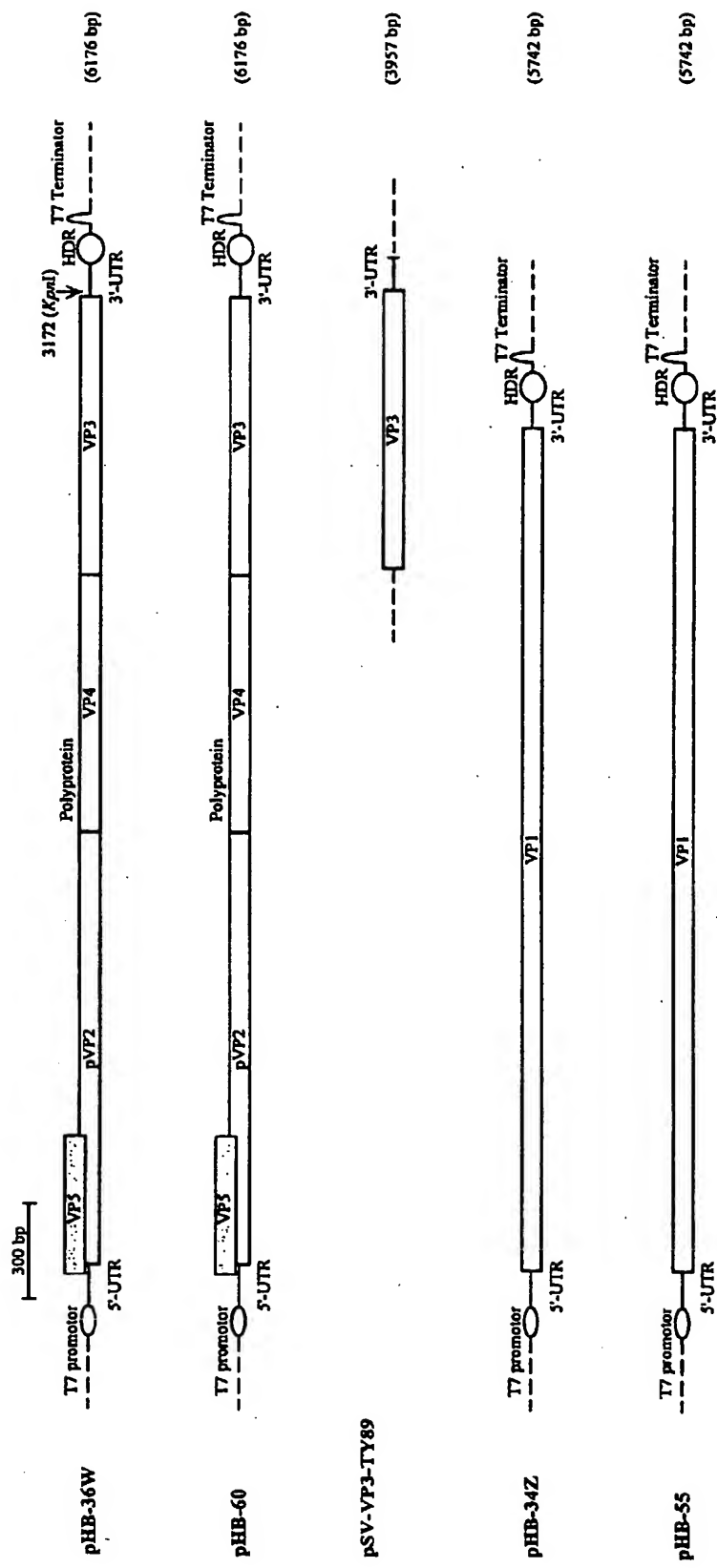
Fig. 3b IBDV VP1 alignment

Consensus	MSD.FNSPQAA RS.ISAAGFI KPTAGQDVER LLIPKVVWPP EDPLASPSRL AKFLRENGYK .LQPRSLPEN KEYET	75
CEF94-VP1	....I.....T.....	75
D6948-VP1	....V.....K.....	75
Consensus	DQILPDLAWH RQIEGAVLKP TLSLPIGDQE YPPKYYPTHR PSKEKPNAYP PDIALLKQMI YLFLQVPEA. .LKD	150
CEF94-VP1	.....	150
D6948-VP1	.....	150
Consensus	EVTLTQNIR DKAYGSGTYM GQATRLVAMK EVATGRNPNK DPLKLGYTFE SIAQLLDITL PVGPPGEDDK PWVPL	225
CEF94-VP1	.....	225
D6948-VP1	.....	225
Consensus	TRVPSRMLVL TGDVVG. PEV EDYLPKINLK SSSGLPVYVGR TKGETIGEMI AISNQFLREL S.LLKQGAGT EGSNK	300
CEF94-VP1	.....D.....	300
D6948-VP1	.....E.....	300
Consensus	KKLLSMLSDY WYLSGCCLLFP KAERYDKRSTW LTKTRNIWSA PSPTHLMISM ITWPVNNSNP NNVLNIEGCP SLYKF	375
CEF94-VP1	.....	375
D6948-VP1	.....	375
Consensus	NPFRGGILNRI VEWI.AP.EP KALVYADNY IVHSNTWYSI DLEKGEANCT RQHMQAAMYY ILTRGWSING DPMFN	450
CEF94-VP1	.....L.E.....	450
D6948-VP1	.....M.D.....	450
Consensus	QTWATTPAMNI APALVVDSSC LIMNLQINTY GQQSGNAAATP INNHLLSTLV LDQMNLM.QP .PDSEEPKSI EDKLG	525
CEF94-VP1	.....	525
D6948-VP1	.....	525
Consensus	INFKIERSID DIRGKILRQIV .IAQPGYLSG GVEPEQ.SPT VELDILLGNSA TYSKDLGTYV FVLDKERLPC SAAYP	600
CEF94-VP1	.....L.....S.....	600
D6948-VP1	.....P.....P.....	600
Consensus	KGVENKSLKS KVIGIEQAYKV VRYEARLRLVG GNNYPLLNKA CKNNA.AARR HLEAKGFPID EPLAENSELS EPGEA	675
CEF94-VP1	.....	675
D6948-VP1	.....	675
Consensus	FEGFNIKLTW T.ESLAELN. PVPPKPPNVN RPVNTGGLKA VSNALRTGRY RNEAGLSGLV LLATARSRLQ DAVKA	750
CEF94-VP1	.....S.....K.....	750
D6948-VP1	.....P.....R.....	750
Consensus	KAEAELHKS KPDDPDADWP ERSETLSDLR EKADIASKVA HSALVETSVA LEAVQSTSYY TPKYPEVNP QTASN	825
CEF94-VP1	.....	825
D6948-VP1	.....	825
Consensus	PVVGHLPLAK RATGVQAALL GAGTSRPMGN EAPTRSKNAV KMAKRRQRQK ESRQ..	881
CEF94-VP1	.....	881
D6948-VP1	.....	879

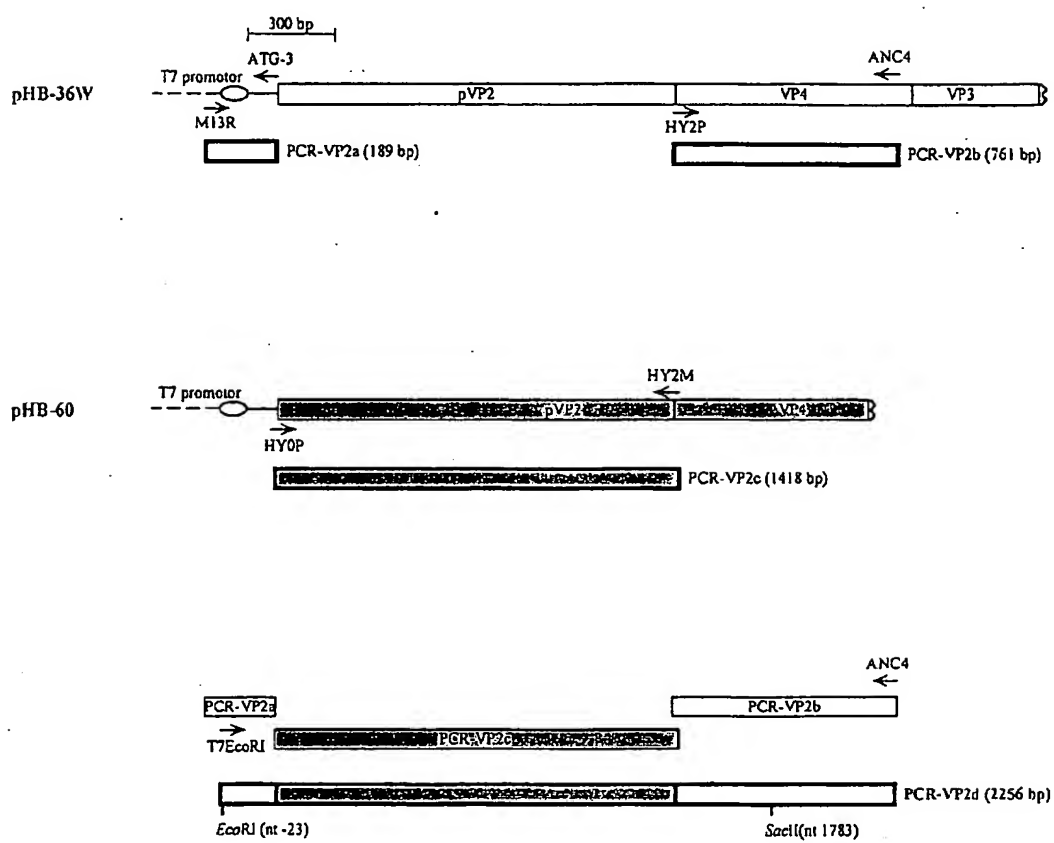
Fig. 3c IBDV VP5 alignment

Consensus	MVS RDQT NDR SDD. PARSNP TDC SVHTEPS DANN RTGVHS GRHP. EAHSQ	50
D6948-VP5	..... E..... .... R.....	50
CEF94-VP5	..... K..... .... G.....	50
Consensus	VRD L DLQ FDC GGH RV RAN CL FPW. PWL NCG CSL H TAE QWE LQ VRS DAP DC	100
D6948-VP5	..... F..... ....	100
CEF94-VP5	..... I..... ....	100
Consensus	PEPT GQL QLL QASE SESHSE VKHT. WWRLC TK. HHK RRDL PRKPE	145
D6948-VP5	..... P..... W.....	145
CEF94-VP5	..... S..... R.....	145

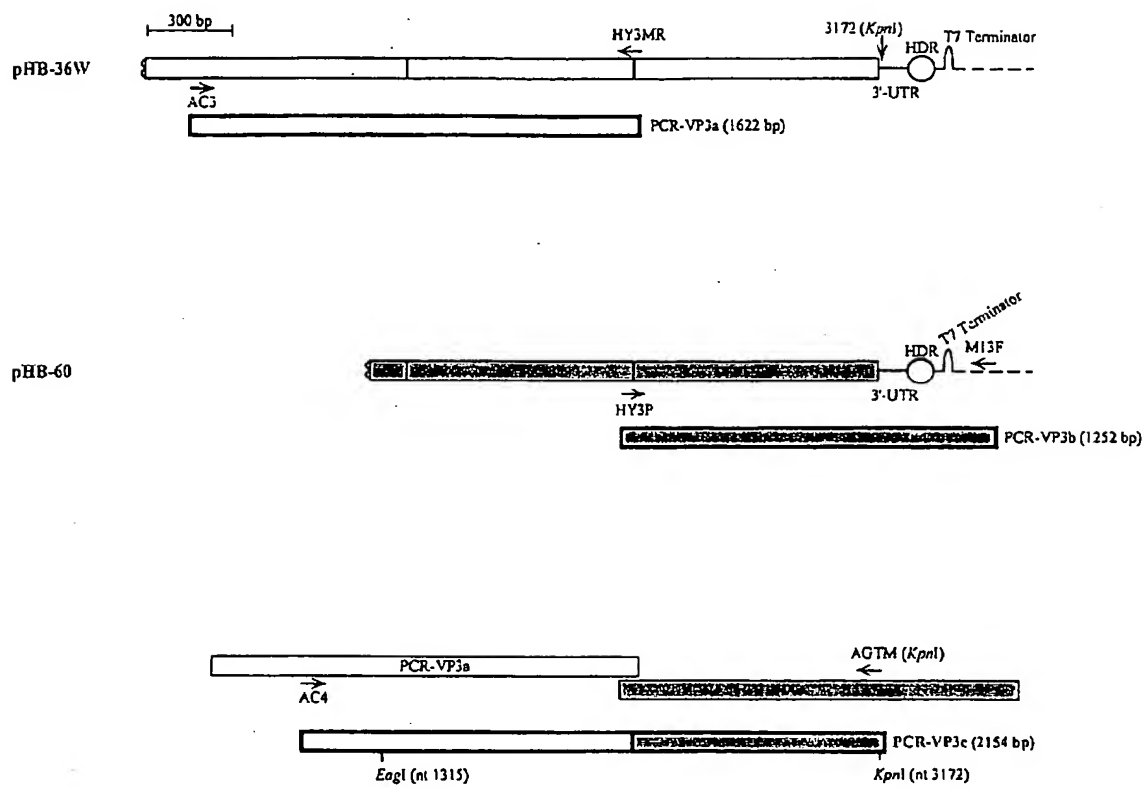
**Fig. 4** Schematic representation of the used plasmids



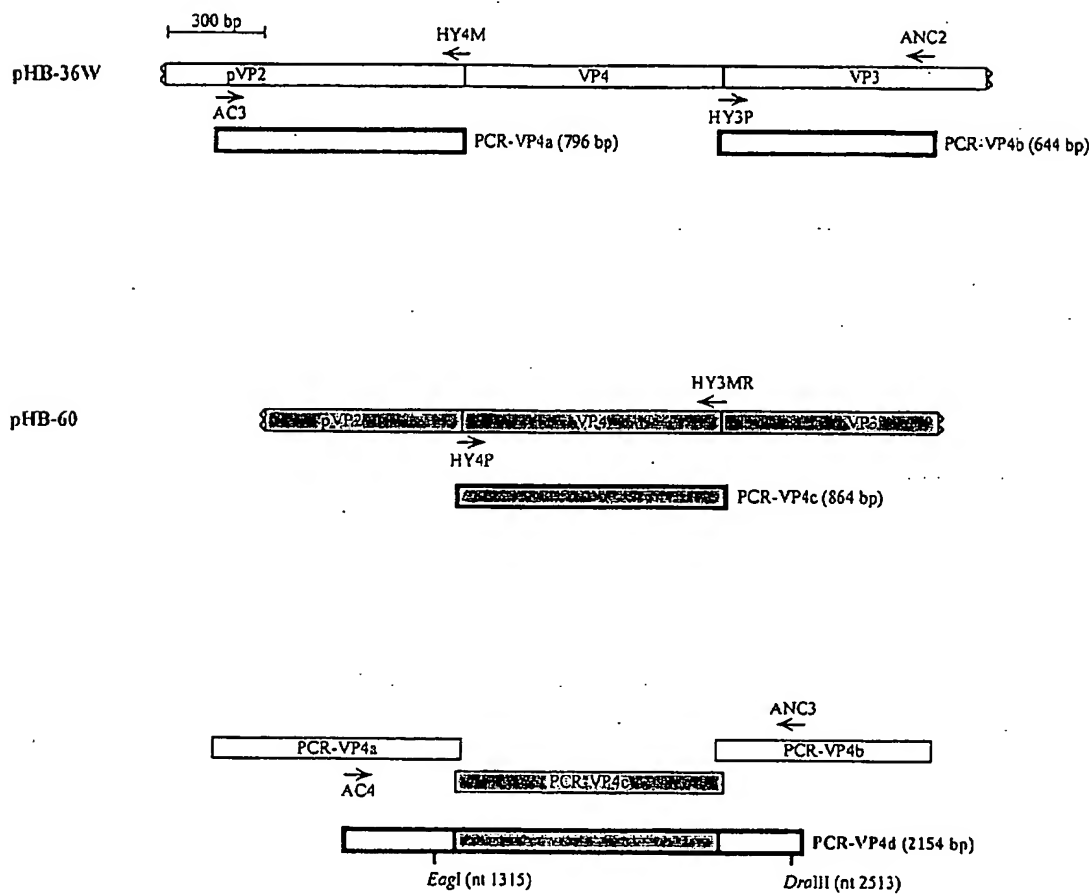
**Fig. 5a** Schematic representation of the construction of PCR fragment PCR-VP2d



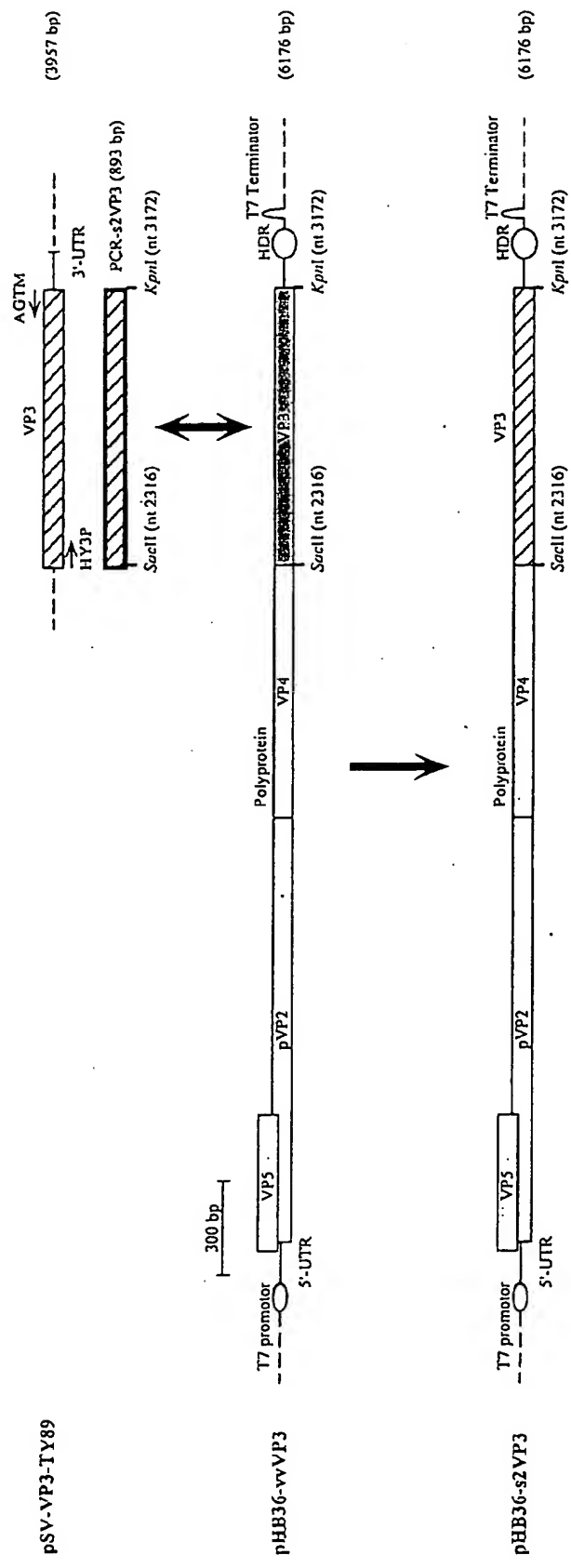
**Fig. 5b** Schematic representation of the construction of PCR fragment PCR-VP3c



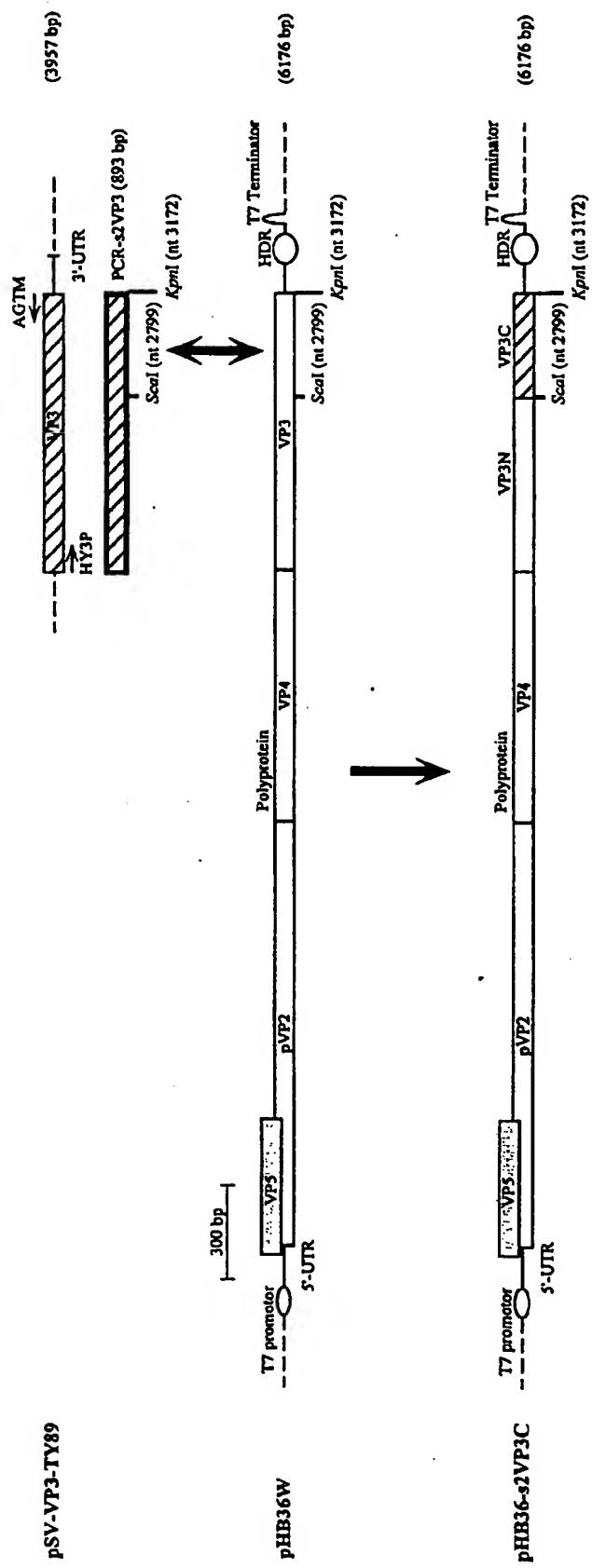
**Fig. 5c** Schematic representation of the construction of PCR fragment PCR-VP4d



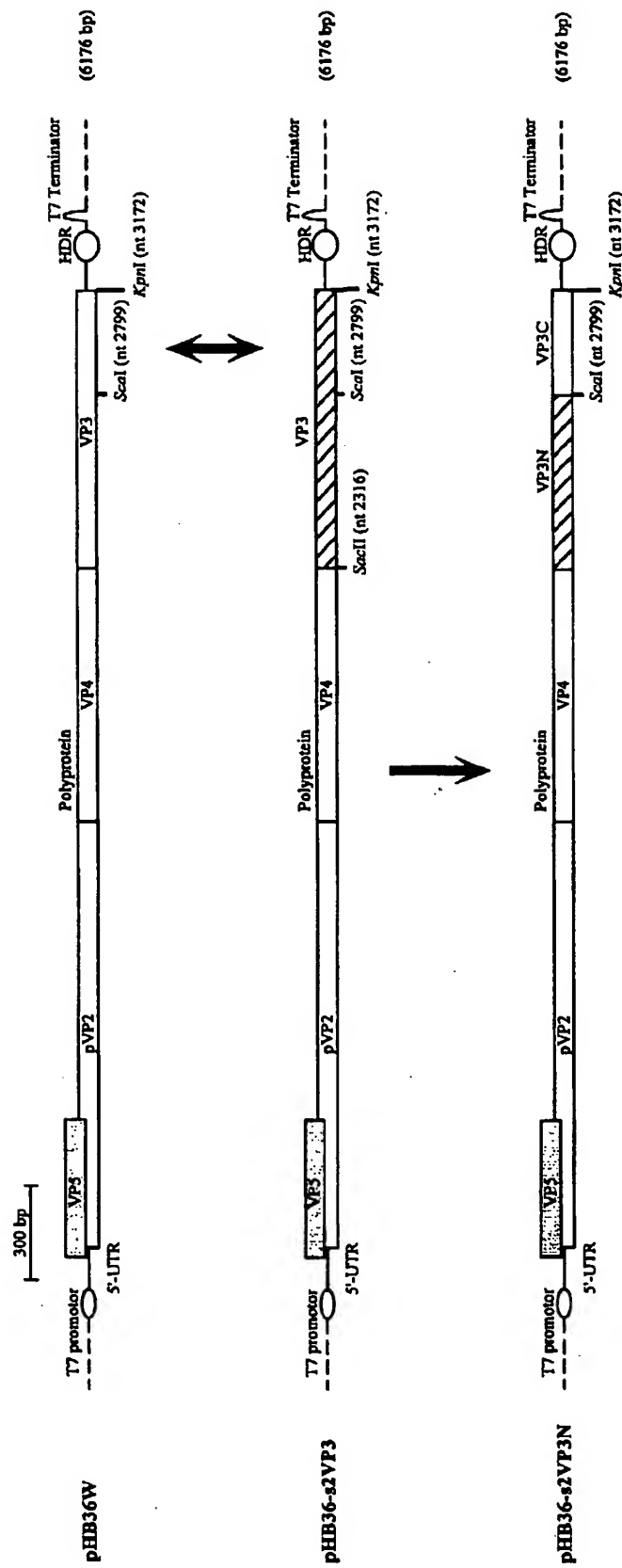
**Fig. 5d** Schematic representation of the construction of plasmid pHB36-s2VP3



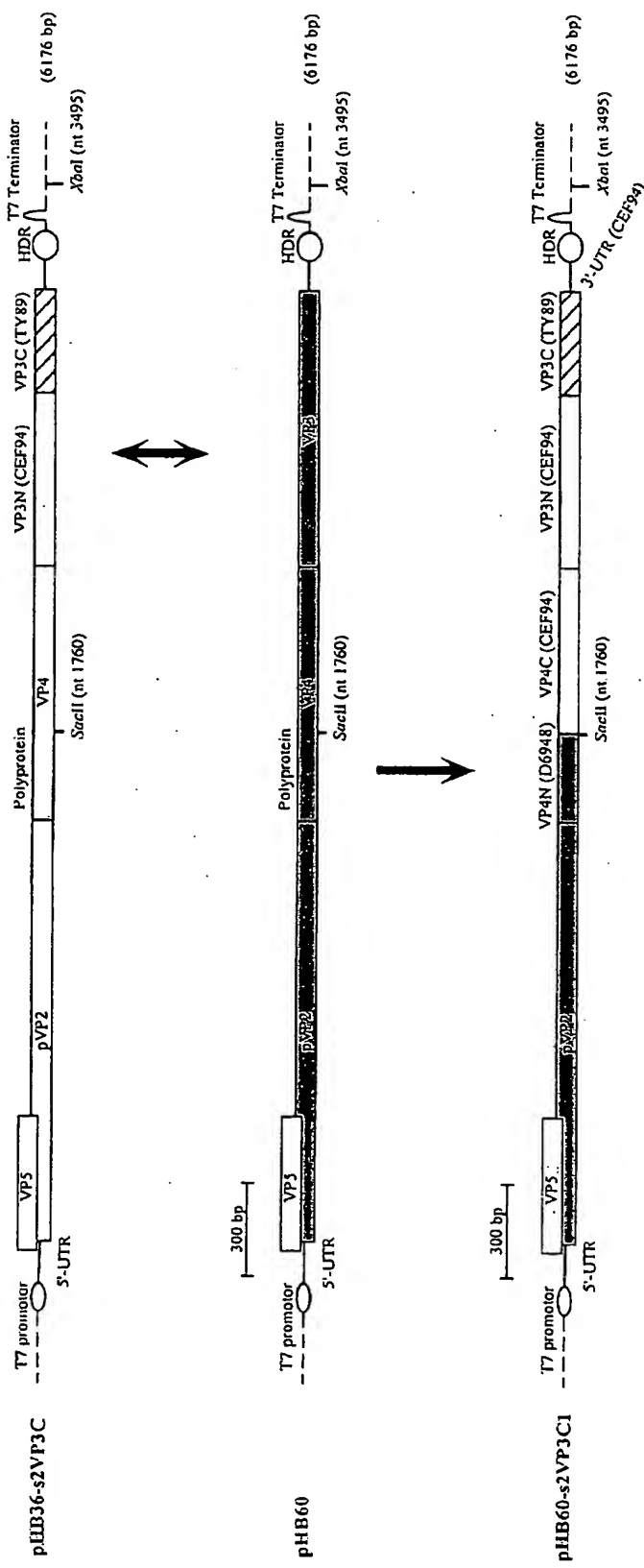
**Fig. 5e** Schematic representation of the construction of plasmid pHB36-s2VP3C



**Fig. 5f** Schematic representation of the construction of plasmid pHB36-s2VP3N



**Fig. 5g** Schematic representation of the construction of plasmid pHB60-s2VP3C1





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## EUROPEAN SEARCH REPORT

Application Number

EP 99 20 2316

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X	WO 98 09646 A (MUNDT EGBERT ;VAKHARIA VIKRAM N (US); UNIV MARYLAND BIOTECHNOLOGY) 12 March 1998 (1998-03-12) * see whole document, particularly claims and page 15, line 18 - page 17 *	1-9, 23-29	
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Place of search	Date of completion of the search	Examiner	
THE HAGUE	6 January 2000	Smalt, R	
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X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document		T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document	

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